• #33/dec.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Atty. Docket: WALLACH=23

In re Application of:

David WALLACH et al

Appln. No.: 09/380,546

Appln. No.: 09/380,546

Filed: November 29, 1999

Washington, D.C.

For: CASH (CASPASE HOMOLOGUE)

WITH DEATH EFFECTOR DOMAIN,)

MODULATORS OF THE FUNCTION

OF FAS RECEPTORS

Art Unit: 1635

Examiner: B. Whiteman

Washington, D.C.

DECLARATION UNDER 37 CFR §1.131

Honorable Commissioner for Patents Washington, D.C. 20231

Sir:

I, Yura "Yury" GOLTSEV, hereby declare and state as
follows:

I am the same Yura Goltsev named as a co-inventor in the above-identified application and my educational and professional experience is presented in the curriculum vitae attached hereto as Exhibit A.

The experiments and computer searches described below were either conducted by me or under my supervision, and I can attest of my own personal knowledge that all the results reported herein and the dates ascribed thereto are true and accurate.

It is my understanding that the claims of the above-identified application remain rejected under 35 U.S.C. \$102(e) as being anticipated by Shu et al., U.S. Patent No. 6,242,569. The filing date of U.S. Patent No. 6,242,569 for purposes of \$102(e) is February 5, 1997.

The present declaration is intended to establish that the invention of the subject matter of the rejected claims was made on or before the effective filing date of U.S. Patent No. 6,242,569. The following shows conception on or before February 5, 1997, and diligence until the filing of Israeli patent application IL 120367, from which the benefit of foreign priority is claimed in the above-identified application, on March 3, 1997, as a constructive reduction to practice.

The dates of the experiments and computer searches to show conception are excised but every excised date is on or before February 5, 1997. Furthermore, the experiments and computer searches were done in Israel, a WTO member country, on or after January 1, 1996.

Attached hereto as Exhibit B are pages 106, 107 and 110 of my laboratory notebook (with the dates excised) with entries made by me in my own handwriting. Page 106 shows a two hybrid (2H) screen of MCH4L with a Jurkat library (in GAD1318 vector) without 3AA and restriction enzyme digests of

clones obtained from the Jurkat library. A continuation of the restriction enzyme digests of the clones from the Jurkat library is shown on page 107. The heading for page 110 is "Cloning of G1 isoforms", where the clones were obtained from RT-PCR of HeLa pA RNA. Restriction enzyme digestions of the clones are shown on page 110, where the "circled" clones were sequenced.

To clarify the terminology used for the polypeptide of the present invention, the Gl α isoform (long splice variant) of SEQ ID NO:2, encoded by the nucleotide sequence of SEQ ID NO:1, is also referred to as CASH α or CASHa, as will be seen in some Exhibits presented herein. Likewise, the Gl β isoform (short splice variant) of SEQ ID NO:4, encoded by the nucleotide sequence of SEQ ID NO:3, is also referred to as CASH β or CASH β .

Two nucleotide sequences, designated j13.1789 and j15.1789, obtained from the G1 isoform clones are shown in Exhibit C. These two nucleotide sequences overlap (but on different strands) to give the nucleotide sequence of the short variant G1 β isoform (CASH β or CASH δ).

For purposes of showing the relationship of the sequences j13.1789 and j15.1789 to CASHb, a color schematic diagram, which is not intended to be to scale, is presented as Exhibit D, where each color represents regions of sequence

identity. The vertical lines starting from the left indicate the start of the coding sequence for CASHa and CASHb, the end of the region of similarity between CASHa and CASHb, the end (bSTOP) of the coding sequence of CASHb, and the end (aSTOP) of the coding sequence of CASHa. As can be seen from this schematic diagram, the nucleotide sequences of j13.1789 and j15.1789 cover the entire coding sequence of CASHb. The sequences labeled with accession numbers AA149562, AA151642, N94588, and W23795 are contigs identified for CASHa and CASHb that were available in the GenBank (NCBI) sequence database prior to the date the invention was made. The sequence database printouts for these contigs are presented as Exhibit E.

Part of the sequence from j15.1789 was used in a BLAST search for identification of similar protein coding regions in the GenBank database. The BLAST search results are presented as Exhibit F and the 313 base pair query sequence used in the BLAST search is shown in Exhibit G.

To search for other splice variants (which is routinely done in the art) of CASHb, a 219 base pair query sequence from j13.1789, shown as Exhibit H, was used in a BLAST search (Exhibit I) for identifying from the GenBank EST database, ESTs with nearly identical sequences. As shown in Exhibit I, the EST sequences with accession numbers AA149562

and AA15142 (highlighted) were identified. The yellow highlighting is not part of the original document and was made contemporaneously herewith to aid the examiner's understanding of this document. EST accession numbers AA149562 and AA151642 are the same accession numbers shown in Exhibit D to generate contigs for the coding sequence of CASHa. Accordingly, from the j15.1789 and j13.1789 sequences generated from the G1 isoform clones and the contigs identified based on BLAST sequence similarity searches, as summarized in Exhibit D, the nucleotide sequences and the encoded amino acid sequences for CASHa and CASHb were obtained, and therefore the present invention was conceived on or before February 5, 1997.

A printout from the "Sequencher" program dated February 8, 1997, supplemented with my hand made drawings on the printout, is attached hereto as Exhibit J. This printout is a contig of ESTs for CASHa and CASHb which lists more ESTs as well as contigs from The Institute for Genomics Research (TIGR) database that have regions coinciding with the CASH sequences.

As the sequences of both the short and long splice variants of CASH (CASHb and CASHa) were obtained by a combination of overlapping clones and contigs identified from a search of an EST sequences database, I set out to confirm the sequence for both CASHb and CASHa by ordering the

synthesis of oligonucleotide primers for use in sequencing/PCR. An ordering form with the heading "Oligos for G1" dated February 9, 1997, and showing the oligo primers ordered, with my handwriting on the side to indicate the identifying number for each the primers ordered, is attached hereto as Exhibit K.

Exhibit L is an updated version of Exhibit J. While it is undated, it was attached to the ordering form dated February 9, 1997, herein presented as Exhibit K. At the bottom of Exhibit L, I had noted in my own handwriting the approximate positions of the oligo primers, i.e., 18046, 18044, etc., first identified in Exhibit K.

Pages 111-113 from my laboratory notebook are attached hereto as Exhibit M, where page 111 shows an analysis of cDNA clones containing CASH sequences recovered from a library and page 112 shows restriction enzyme digests of plasmids containing cDNA fragments subcloned from the clones presented on page 111. While page 111 is undated, I performed this experiment either before or contemporaneously with the experiment on page 112. Page 113, also undated, shows a restriction enzyme analysis of plasmids expressing deletion mutants of CASH.

Attached hereto as Exhibit N is the printout from a BLAST search of the GenBank EST database using a 139 base pair

sequence from murine CASH as the query sequence. Note that on page 3 of this printout the BLAST search was conducted on February 16, 1997.

The sequences identified by 10609.1818, 11717.1818, and 18648.1818, dated February 25, 1997, and attached hereto as Exhibit O, are from sequencing a CASHb expression vector using three different primers, one of which (primer 18648) is shown on the oligo ordering form presented as Exhibit K.

Similarly, sequencing of a CASHa expression vector yielded the sequences identified by rti-651.1831 and rt1-648.1831, dated March 3, 1997, and attached hereto as Exhibit P.

The sequences of some "V" clones containing CASH (both long and short) obtained from cDNA library screening, dated February 25, 1997, are presented in Exhibit Q.

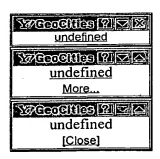
Exhibit R, attached hereto, presents three sheets that were attached together in my datafile. The first page is an undated schematic hand made drawing of V clones containing CASH (both long and short). The second page is a "Sequencher" contig, dated February 26, 1997, of ESTs and the sequences of some the V clones. The third page, dated February 27, 1997, is a more expanded contig including ESTs and the sequences for V clones.

In conclusion, the conception of the present invention occurred on or before February 5, 1997, and there was diligence from February 5, 1997, the date of filing of U.S. Patent No. 6,242,569, until the March 3, 1997, filing of Israeli patent application IL 120367 as a constructive reduction to practice. Accordingly, the applied Shu et al. reference, U.S. Patent No. 6,242,569, is antedated and withdrawal of the 35 U.S.C. §102(e) rejection is warranted.

The undersigned declares further that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

27 may 2003

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CURRICULUM VITAE

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EDUCATION:

Bachelor of Science		
Moscow Institute for Physics and		
Technology Majors in Physical		
Chemistry and Biophysics		
Master of Science		
Moscow Institute for Physics and		
Technology		
Subject area: Molecular Biophysics		
Ph.D. in Molecular Biology		
The Weizmann Institute of Science		
Supervisor: Prof. David Wallach		
Thesis topic: Novel approaches to		
identification of signaling molecules		
activated by receptors of the TNF/NGF		
family, and their application to the study		
of cFLIP/CASH		

DATE OF BIRTH: June 10th, 1972

MARITAL STATUS: married, two children

CITIZENSHIP: Israel

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AREA OF SCIENTIFIC EXPERTISE:

Last six years: Molecular analysis of cytokine-induced signal transduction pathways, in particular those triggered by the TNF/FAS-family members. Molecular regulation of apoptosis.

TECHNICAL SKILLS:

Recombinant DNA: conventional molecular cloning techniques, RT-PCR, site-directed mutagenesis, construction of cDNA libraries, DNA & RNA expression analysis (electrophoresis, hybridization, protection, etc), library screening, subtraction by Representation Difference Analysis (RDA)

Mammalian Cell Culture: routine, expression of recombinant protein in culture cells (both transient & stable), analysis: Western, Far Western, immunocytochemistry, GFP visualization system, essential flow cytometry and sorting, ELISA.

Signaling-related Techniques: EMSA & reporter assays for transcription, *in vitro* kinase assay (in gel & in solution), cell viability assays (biochemical and visual), *in vivo* labeling, *in vivo* signal transduction pathway reporting systems (e.g. PathFinder from Stratagene).

Protein-protein Interactions Assays: yeast two- and three-hybrid systems; mammalian interaction traps, *in vitro* pull-down, *in vivo* co-immunoprecipitation (both endogenous and transfected proteins).

Proteomics: Tandem Affinity Purification (TAP) of protein complexes and identification of the components by microsequencing.

Bacterial Expression: Introduction and affinity purification of recombinant proteins, essential HPLC (gel filtration, ion exchange)

PUBLICATIONS:

Boldin, M. P., Goncharov, T. M., Goltsev, Y. V., and Wallach, D. (1996). Involvement of MACH, a novel MORT1/FADD-interacting protease, in Fas/APO-1- and TNF receptor-induced cell death, Cell 85, 803-15.

Goltsev, Y. V., Kovalenko, A. V., Arnold, E., Varfolomeev, E. E., Brodianskii, V. M., and Wallach, D. (1997). CASH, a novel caspase homologue with death effector domains, J Biol Chem *272*, 19641-4.

Wallach, D., Boldin, M., Goncharov, T., Goltsev, Y., Mett, I., Malinin, N., Adar, R., Kovalenko, A., and Varfolomeev, E. (1996). Exploring cell death mechanisms by analyzing signaling cascades of the TNF/NGF receptor family, Behring Inst Mitt, 144-55.

Wallach, D., Varfolomeev, E. E., Malinin, N. L., Goltsev, Y. V., Kovalenko, A. V., and Boldin, M. P. (1999). Tumor necrosis factor a receptor and Fas signaling mechanisms, Annu Rev Immunol 17, 331-67.

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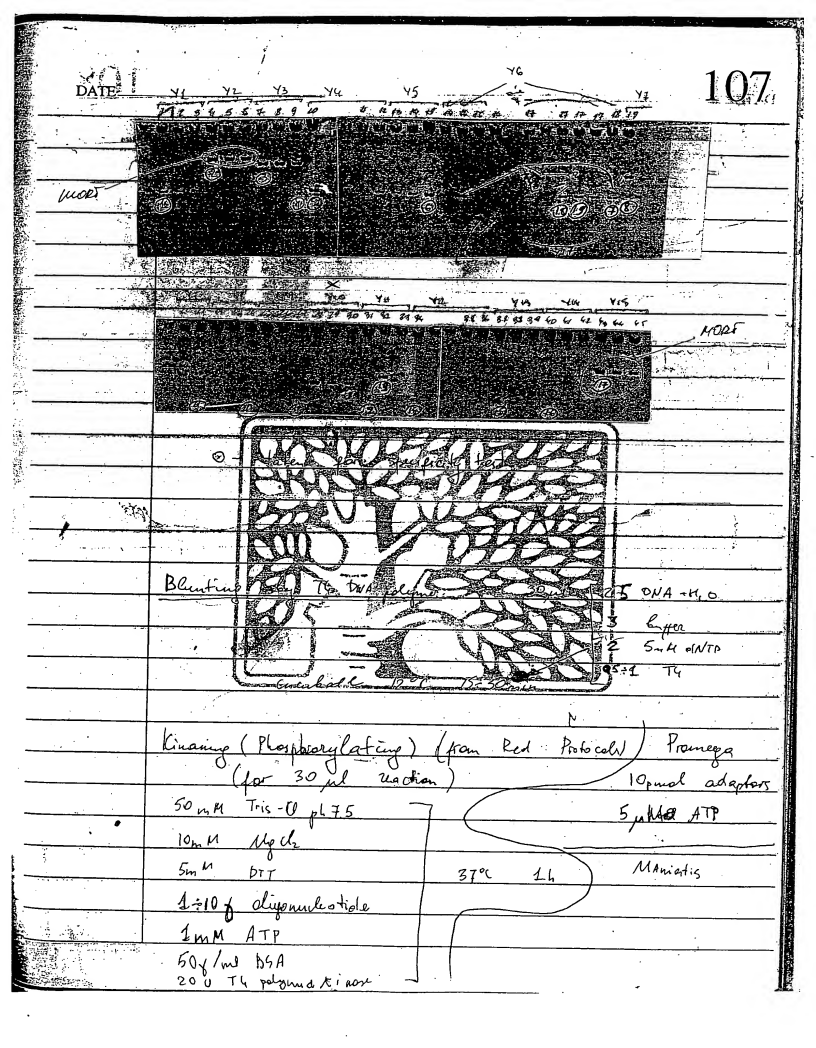
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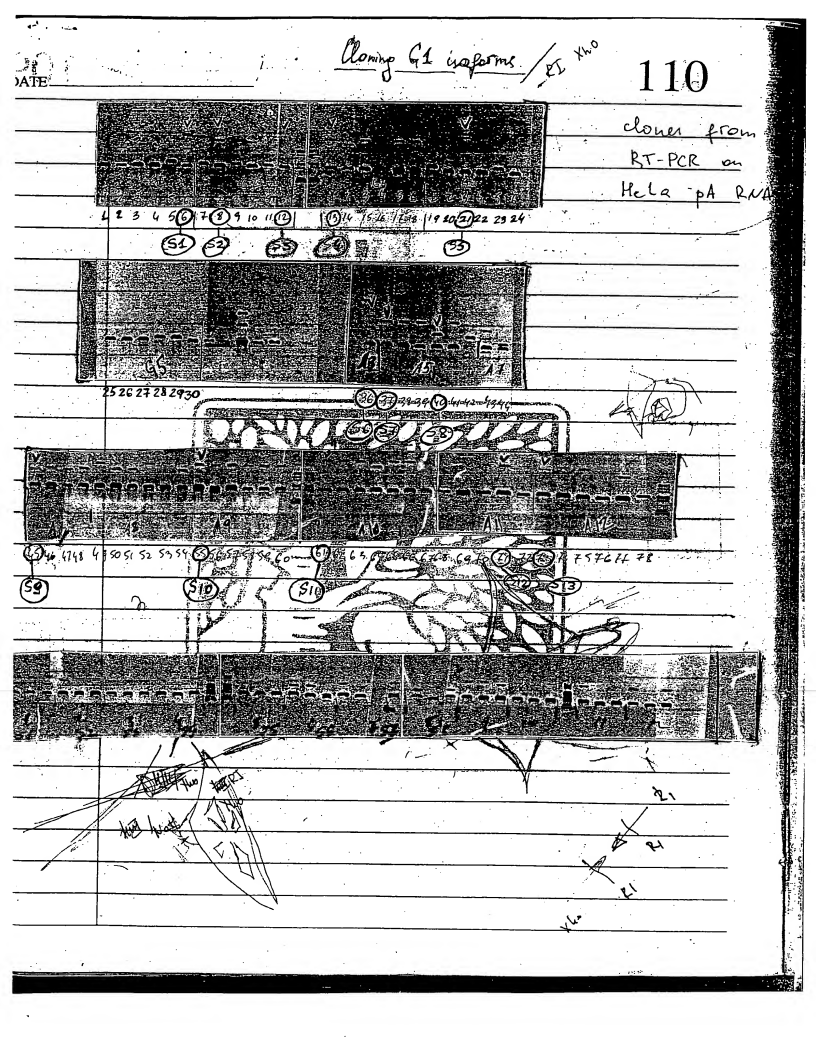
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- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

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our Ref: SampleB-1789 27 Type: N Check: 3143 CGATTCATAG ATCAGGGTTT TCCCAGTCAC GACGTTGTAA AACGACGGCC AGTGGAATTG TAATACGACT CACTATAGGG CGAATTGGGT ACCGGGCCCC 51 CCCTCGAGGT CGACCGCCG CAGGGAATTT TTTTTTTTT TTTTTTTTT 101 TTTTTTTTT AAAAAGTACA AGGGATATTT ATTTTATAGC ATTATTAGTA 151 GTAACAAAG ATTGGAAAGC ATCTAAACAT ACAGCAACAA GAAAGGGCTa 201 AACACATTAT AACACATTTA GAAAATGAAA CGATTTAATC ATTCCCAATG 251 ATAAANTTNA GGCATTCCAT NTTNNNCACA TGGAANAATT TCCANNCAAT 301 351 TTTCNTATCA TGACAGGGGG CTTNGGGTGT TNTCTNCCTN AANTTTTTTN 401 AAGGATCCTT NAAACTCTTT TGGATTGCTG CTTGGANAAA NTTCCTGTAA CTTGTCCCTG CNCCTATNAA NNGANTGCAG GGTACTTCTG GATTTTTGTC 451 501 TTCAGGTCTA TTCTGTGGAT GTTCTTTAGG CATTTTTCTA ATAAATCCCT TGATCTGGGG CCCCCAATTT ANTTTCTCCC ACTCAACCAC AAGGTCCAAA 551 601 AAACTCTTCT CCNNGCTTAT CTTGCCTCGG CCCCTGTTTC CTTCCTTGAN 651 GAAAATTATN ANGACNCTCC CATTTTCCNA NTCCTCCCCA TCTCTGCCAT 701 CACCCTCTAT TTCCNAAACA AGGTGANGGT TCTTACCAGG TGGGTTCCCN 751 CNCNTTTCTN TCCTCTTCNA AANACTTTGA NCGGTCCNNT CCCCCCCNTT 801 TAANATNCAC CANTCCCCAC CAAAAATTTC TCTTTNCCNT CAATTCCAAA 851 GTCCNGNCTT AGGTGACCCA CTTTCCANCN CCGGCCNNAA AANACNCCCC 901 CCCCCNTTTA GTTCCCTCCC CTGAAGGTN

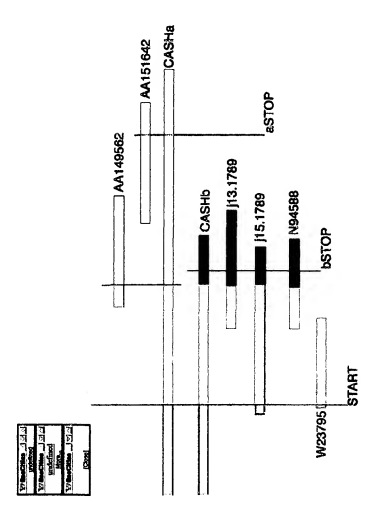
Exhibit C

your ID: j15.1789 j15.1789.seq Lengt. our Ref: SampleB-1789 22 Check: 7805 CCCACCAAAC CAAAAAAAGA GNCCTAGAAC TAGTCGCAAATTAGGGCACGAG CTGTCTGCTG AAGTCATCCA TCAGGTTGAA GAAGCACTTG ATACAGATGA GAAGGAGATG CTGCTCTTTT TGTGCCGGGA TGTTGCTATA GATGTGGTTC 101 CACCTAATGT CAGGGGACCT TCTGGATATT TTACGGGAAA GAGGTAAGCT 151 GTCTGTCGGG GACTTGGCTG AACTGCTCTA CAGAGTGAGG CGATTTGACC 201 TGCTCAAACG TATCTTGAAG ATGGACAGAA AAGCTGTGGA GACCCACCTG 251 CTCAGGAACC CTCACCTTGT TTCGGACTAT AGAGTGCTGA TGGCAGAGAT 301 TGGTGAGGAT TTGGATAAAT CTGATGTGTC CTCATTAATT TTCCTCNTGA 351 AGGATTACAT GGGCCGAGGC AANATAATCA AGGAGAAGAN TTTCTTGGAC 401 CTTGTGGTTG AATTGGGAGA AACTAAATCT GGTTGCCCCN CNATCAACTG 451 GATTTATTAT AAAAATGCCT AAAGAACTCC CCCCANTANA CCTGAAGACA 501 AAAATCCANA ANTACTTCTN TCTGTNCNCG GANCANGGAC GANTTNCTNG 551 AATGTTCTCC TANCANCATC CNTNAGAATC TCCANGATCC TTCCNNTNAC 601 TTCNNGATGA TACCCCTNTC CCCNTGTCCT NATCTGAAAT TCTGGAAATT 651 GTCCATNTAA TTACNTGGAC TGCCTCNCNT TATCTTCNGN AACNAATAAA 7-01 TCTTCCCTNC CTAATGTTTT TNGAGNNTTT ACCCTCCCCG TTNCTGTTTN 751 TTANATCCTC CCCCCCTTG TTTCTCTNTT ATTCNNNNAA ANANTTCCCT 801 TCTTTCTTTT NNNNCCCCCN NNNNNATAA NAATNNTCCC CCCCACNCAG 851

GGGGNNCGGC CCCNTCCCNT TTTAATTT

901

4 777





ATCGGATCCCCGGCAGE (2) EGCTCAGGAT SACTTCG CCCCA C CTTCGCATACGTCTTECNTBLOGICOS CONTESCONOS CO EACAGACT

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Taxonomy

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Nucleotide

Show: 20

Protein

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1: AA149562. zo29e05.rl Strata...[gi:1720363]

Limits

Links

IDENTIFIERS

dbEST Id: EST name: GenBank Acc:

788753 zo29e05.rl AA149562 1720363

GenBank gi: GDB Id:

4620622

CLONE INFO

Clone Id:

IMAGE:588320 (5')

Source:

IMAGE Consortium, LLNL

CDNA DNA type:

PRIMERS

Sequencing:

-28M13 rev2 from Amersham

PolyA Taiĺ:

Unknown

SEQUENCE

GGAGCAGGGACAAGTTACAGGAATGTTCTCCAAGCAGCAATCCAAAAGAGTCTCAAGGAT CCTTCAAATAACTTCAGGCTCCATAATGGGAGAAGTAAAGAACAAAGACTTAAGGAACAG CTTGGCGCTCAACAAGAACCAGTGAAGAAATCCATTCAGGAATCAGAAGCTTTTTTGCCT CAGAGCATACCTGAAGAGATACAAGATGAAGAGCCAAGCCCCTAGGAATCTGCCTGATA ATCGATTGCATTGGCAATGAGACAGAGCTTCTTCGAGACACCTTCACTTCCCTGGGCTAT GAAGTCCAGAAATTCTTGCATCTCAGTATGCATGGTATATCCCAGATTCTTGGCCAATTT GGTCCCAGAGTGTGTATGGTGTGGATCAGACTCACTCAGGGCTCCCCTGCATCACATCAG GAGGATGTTCATGGGAGAATCATGCCCTTATCTAGCAGGAAGCAAAGATGTTTTTATC

Quality:

High quality sequence stops at base: 398

Entry Created:
Last Updated:

Dec 10 1996 Dec 10 1996

COMMENTS

This clone is available royalty-free through LLNL; contact the IMAGE Consortium (info@image.llnl.gov) for further

information.

LIBRARY

Lib Name:

Stratagene colon (#937204)

Organism:

Homo sapiens

Organ: Tissue type: colon

tumor

Cell line:

T84 carcinoma cell line

Lab host:

SOLR cells (kanamycin resistant)

Vector:

pBluescript SK-

R. Site 1: R. Site 2: **ECORI**

XhoI

Description:

Cloned unidirectionally. Primer: Oligo dT. T-84 colonic epithelial cell line. Average insert size: 1.0 kb; Uni-ZAP XR Vector; ~5' adaptor sequence: 5' GAATTCGGCACGAG 3' ~3'

adaptor sequence: 5' CTCGAGTTTTTTTTTTTTTTT 3'

SUBMITTER

Name:

Wilson RK

Institution:

Washington University School of Medicine

Address:

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Tel: Fax: 314 286 1800 314 286 1810

E-mail:

est@watson.wustl.edu

CITATIONS

Medline UID:

97044478

Title:

Generation and analysis of 280,000 human expressed sequence

tags

Authors:

Hillier, L., Lennon, G., Becker, M., Bonaldo, M.F., Chiapelli, B., Chissoe, S., Dietrich, N., DuBuque, T., Favello, A., Gish, W., Hawkins, M., Hultman, M., Kucaba, T., Lacy, M., Le, M., Le, N., Mardis, E., Moore, B., Morris, M., Parsons, J., Prange, C., Rifkin, L., Rohlfing, T., Schellenberg, K., Soares, M.B., Tan, F., Thierry-Meg, J., Trevaskis, E., Underwood, K., Wohldmann, P.,

Waterston, R., Wilson, R., Marra, M. Genome Res. 6 (9): 807-828 1996

Citation:

MAP DATA

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Structure

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Taxonomy

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Genome

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☐ 1: AA149562. zo29e05.r1 Strata...[gi:1720363]

Links

Details

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IDENTIFIERS

dbEST Id:

788753

EST name: GenBank Acc:

zo29e05.r1 AA149562 1720363

GenBank gi: GDB Id:

4620622

CLONE INFO

Clone Id:

IMAGE:588320 (5')

Source:

IMAGE Consortium, LLNL

DNA type:

CDNA

PRIMERS

Sequencing:

-28M13 rev2 from Amersham

PolyA Tail:

Unknown

SEQUENCE

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Quality:

High quality sequence stops at base: 398

Entry Created: Last Updated:

Dec 10 1996 Dec 10 1996

COMMENTS

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the IMAGE Consortium (info@image.llnl.gov) for further

information.

LIBRARY

Lib Name:

Stratagene colon (#937204)

Organism:

Homo sapiens

Organ:

colon tumor

Tissue type: Cell line:

T84 carcinoma cell line

Lab host:

SOLR cells (kanamycin resistant)

Vector:

pBluescript SK-

R. Site 1:

EcoRI

R. Site 2:

XhoT

Description:

Cloned unidirectionally. Primer: Oligo dT. T-84 colonic epithelial cell line. Average insert size: 1.0 kb; Uni-ZAP XR Vector; ~5' adaptor sequence: 5' GAATTCGGCACGAG 3' ~3'

adaptor sequence: 5' CTCGAGTTTTTTTTTTTTTT 3'

SUBMITTER

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CITATIONS

Medline UID:

97044478

Title:

Generation and analysis of 280,000 human expressed sequence

tags

Authors:

Hillier, L., Lennon, G., Becker, M., Bonaldo, M.F., Chiapelli, B., Chissoe, S., Dietrich, N., DuBuque, T., Favello, A., Gish, W., Hawkins, M., Hultman, M., Kucaba, T., Lacy, M., Le, M., Le, N., Mardis, E., Moore, B., Morris, M., Parsons, J., Prange, C.,

Rifkin, L., Rohlfing, T., Schellenberg, K., Soares, M.B., Tan, F., Thierry-Meg, J., Trevaskis, E., Underwood, K., Wohldmann, P.,

Waterston, R., Wilson, R., Marra, M.

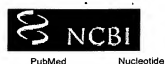
Citation:

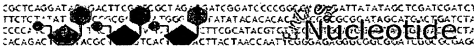
Genome Res. 6 (9): 807-828 1996

MAP DATA

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Apr 28 2003 10:17:55





Search Nucleotide for Genome Structure **PMC**

Taxonomy Cient. **OMIM** Boo

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Show: 20 Q

Protein

° ad banes '

File ∇

□1: N94588. zb79a05.s1 Soares...[gi:1266897]

Links

IDENTIFIERS

dbEST Id:

501376

EST name: GenBank Acc:

zb79a05.s1 N94588

GenBank gi: GDB Id:

1266897 1253190

CLONE INFO

Clone Id: IMAGE: 309776 (3')

Source:

IMAGE Consortium, LLNL

Insert length:

939 **cDNA**

DNA type: PRIMERS

Sequencing:

mob.REGA+ET

PolyA Tail:

Unknown

SEQUENCE

GAAGTACAAGGATATTTATTTTATAGCATTATTAGTAGTAACAAAAGATTGGAAAGCATC TAAACATACAGCAACAAGAAAGGGCTAAACACATTATAACACATTTAGAAAATGAAACGA TTTAATCATTCAGAATGATTAAGTAGAGGCAGTTCCATGTTAATCACATGGAACAATTTC CAAGAATTTTCAGATCAGGACAATGGGCATAGGGTGTTATCATCCTGAAGTTATTTGAAG GATCCTTGAGACTCTTTTGGATTGCTGCTTGGAGAACATTCCTGTAACTTGTCCCTGCTC CTTTGAACAGACTGCTTGTACTTCTGGATTTTTGTCTTTCAGGTCTATTCTGTGGATGTT $\tt CTTTAGGCATTTTCTAATAAATCCAGTTGATCTGGGGCAACCAGATTTAGTTTCTCCAA$

CTCAAC

Quality:

High quality sequence stops at base: 292

Entry Created:

May 9 1996

Last Updated:

Aug 20 1996

COMMENTS

This clone is available royalty-free through LLNL; contact the IMAGE Consortium (info@image.llnl.gov) for further

information.

LIBRARY

Lib Name:

Soares_senescent_fibroblasts_NbHSF

Organism:

Homo sapiens

Tissue type:

senescent fibroblast

Lab host:

DH10B (ampicillin resistant)

Vector:

pT7T3D (Pharmacia) with a modified polylinker V_TYPE:

phagemid

R. Site 1:

Not I

R. Site 2:

Eco RI

Description:

1st strand cDNA was primed with a Not I - oligo(dT) primer

SUBMITTER

Name:

Wilson RK

Institution:

Washington University School of Medicine

Address:

4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108

Tel:

314 286 1800 314 286 1810

Fax: E-mail:

est@watson.wustl.edu

CITATIONS

Medline UID:

97044478

Title:

Generation and analysis of 280,000 human expressed sequence

tags

Authors:

Hillier, L., Lennon, G., Becker, M., Bonaldo, M.F., Chiapelli, B., Chissoe, S., Dietrich, N., DuBuque, T., Favello, A., Gish, W., Hawkins, M., Hultman, M., Kucaba, T., Lacy, M., Le, M., Le, N.,

Mardis, E., Moore, B., Morris, M., Parsons, J., Prange, C.,

Rifkin, L., Rohlfing, T., Schellenberg, K., Soares, M.B., Tan, F., Thierry-Meg, J., Trevaskis, E., Underwood, K., Wohldmann, P.,

Waterston, R., Wilson, R., Marra, M. Genome Res. 6 (9): 807-828 1996

Citation:

MAP DATA

<u>Disclaimer | Write to the Help Desk</u> <u>NCBI | NLM | NIH</u>

Apr 28 2003 10:17:55





Search Nucleotide for

Taxonomy Go Clear Boo

ම්පුව්වූද් default

Limits Show: 20

Preview/Index

History File Clipboard

Details

☐ 1: W23795. zb79a05.rl Soares...[gi:1300619]

Links

IDENTIFIERS

PubMed

dbEST Id: EST name: **530869** zb79a05.rl W23795

GenBank Acc:
GenBank gi:
GDB Id:

1300619 1253190

CLONE INFO

Clone Id:

IMAGE:309776 (5')

Source:

IMAGE Consortium, LLNL

Insert length:
DNA type:

939 CDNA

PRIMERS

Sequencing: PolyA Tail:

mob.REGA+ET Unknown

SEQUENCE

CAGAGATTGGTGA

Quality:

High quality sequence stops at base: 247

Entry Created:

Last Updated:

May 9 1996 Aug 20 1996

COMMENTS

This clone is available royalty-free through LLNL; contact

the IMAGE Consortium (info@image.llnl.gov) for further

information.

LIBRARY

Lib Name:

Soares senescent fibroblasts NbHSF

Organism:

Homo sapiens

Tissue type:

senescent fibroblast

Lab host:

DH10B (ampicillin resistant)

Vector:

pT7T3D (Pharmacia) with a modified polylinker V_TYPE:

phagemid

R. Site 1:

Not I

R. Site 2:

Eco RI

Description:

1st strand cDNA was primed with a Not I - oligo(dT) primer

adapters (Pharmacia), digested with Not I and cloned into the Not I and Eco RI sites of a modified pT7T3 vector

(Pharmacia). Library went through one round of normalization

to a Cot = 5. Library constructed by Bento Soares and M.Fatima Bonaldo.

SUBMITTER

Name:

Wilson RK

Institution:

Washington University School of Medicine

Address:

4444 Forest Park Parkway, Box 8501, St. Louis, MO 63108

Tel: Fax:

314 286 1800 314 286 1810

E-mail:

est@watson.wustl.edu

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Medline UID:

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Title:

Generation and analysis of 280,000 human expressed sequence

Authors:

Hillier, L., Lennon, G., Becker, M., Bonaldo, M.F., Chiapelli, B. , Chissoe, S., Dietrich, N., DuBuque, T., Favello, A., Gish, W., Hawkins, M., Hultman, M., Kucaba, T., Lacy, M., Le, M., Le, N., Mardis, E., Moore, B., Morris, M., Parsons, J., Prange, C.,

Rifkin, L., Rohlfing, T., Schellenberg, K., Soares, M.B., Tan, F.

, Thierry-Meg, J., Trevaskis, E., Underwood, K., Wohldmann, P.,

Waterston, R., Wilson, R., Marra, M. Genome Res. 6 (9): 807-828 1996

Citation:

MAP DATA

<u>Disclaimer</u> | Write to the Help Desk NCBI | NLM | NIH

Apr 28 2003 10:17:55

BLAST Search Results with RepeatMasker filtering, Entrez and SRS links

BLAST search performed using the National Center for Biotechnology Information's <u>BLAST WWW</u> <u>Server</u>. Repeat reporting and filtering performed by <u>RepeatMasker</u> from U. Washington.

Links to Entrez and to the Sequence Retrieval System (SRS) provided by the Human Genome Center, Baylor College of Medicine:

= Retrieve Entrez links (e.g., Medline abstracts, FASTA-formatted sequence reports).

= Retrieve Entrez links to Related sequences (neighbors).

S = Retrieve SRS links (if present).

Reference: Gish, Warren and David J. States (1993). Identification of protein coding regions by database similarity search. Nat. Genet. 3:266-72. Altschul, Stephen F., Warren Gish, Webb Miller, Eugene W. Myers, and David J. Lipman (1990). Basic local alignment search tool. J. Mol. Biol. 215:403-10.

Notice: statistical significance is estimated under the assumption that the equivalent of one entire reading frame in the query sequence codes for protein and that significant alignments will involve only goding reading frames.

Translating both strands of query sequence in all 6 reading frames

Database: Non-redundant GenBank CDS

translations+PDB+SwissProt+SPupdate+PIR

241,922 sequences; 68,723,500 total letters.

Searching.....done

Smallest Sum Reading High Probability Sequences producing High-scoring Segment Pairs: Frame Score gi | 1718327 (U75698) ORF K13 [Kaposi's sarcom... +3 gi|1492103 (U60315) MC160L [Molluscum contag... +3 pir/1855668 hypothetical protein E8 - equine ... +3 4.2e-05 gil1492102 (U60315) MC159L [Molluscum contag... +3 63 9.5e-05 gil1813378 (D86547) still life type 1 [Droso... -3 <u>4.0.</u> 0.00040 Sp/P47211 | GALR HUMAN GALANIN RECEPTOR (GAL1-R) /gi|559... +2 <u>73</u> 0.26 <u>ail58271</u> (X12896) CTAP-III protein (AA 1-8... +3 46 0.59 gi1208127 (M88539) Col-CTAP-III(Leu21) smal... +3 46 2 0.70 gi|1685110 (U58210) tetrahydrofolate dehydro... -3 68 0.79 gi1791038 (X84709) mediator of receptor ind... +3 <u>51</u> _ 0.87 pirllA56912 FADD protein - human /gi|809487 (... +3 <u>51</u> 0.87 gb|1002001 Sequence 1 from Patent US 4897348... +3 44 0.91 gi1208159 (M11517) connective tissue activa... +3 44 0.92 sp1P223661MY88 MOUSE MYELOID DIFFERENTIATION PRIMARY R... +3 0.95 <u>66</u> 43 gi1807578 (M19420) unknown protein [Germist... +1 0.98 splP29425|TX26 PHONI NEUROTOXIN TX2-6 /pir||S29216 neu... +1 34 0.99 splp02775 | pf4L human platelet basic protein precursor ... +3 0.9998

ER

gil1718327 (U75698) ORF K13 [Kaposi's sarcoma-associated herpesvirus]

Length = 139

Plus Strand HSPs:

Score = 121 (55.7 bits), Expect = 3.0e-09, P = 3.0e-09Identities = 31/97 (31%), Positives = 51/97 (52%), Frame = +3

Query: 18 EVIHQVEEALDTDEKEMLLFLCRDVAIDVVPPNVRDLLDILRERGKLSVGDLAELLYRVR 197

EV+ +V L TD++E++LFL + L L+E G+L+ LAE L+R

Sbjct: 5 EVLCEVARKLGTDDREVVLFLLNVFIPQPTLAQLIGALRALKEEGRLTFPLLAECLFRAG 64

Query: 198 RFDLLKRILKMDRKAVETHLLRNPHLVSDYRVLMSEI 308

R DLL+ +L +D + +E HL S Y++ + +

Sbjct: 65 RRDLLRDLLHLDPRFLERHLAGTMSYFSPYQLTVLHV 101

ER

gi!1492103 (U60315) MC160L [Molluscum contagiosum virus subtype 1]

Length = 371

Plus Strand HSPs:

Score = 69 (31.7 bits), Expect = $7.4e^{-06}$, Sum P(2) = $7.4e^{-06}$ Identities = 17/39 (43%), Positives = 24/39 (61%), Frame = +3

Query: 123 DLLDILRERGKLSVGDLAELLYRVRRFDLLKRILKMDRK 239

D L L+ R L++ +AELL +RRFD+LK M R+

Sbjct: 42 DALRALQRRRLLTLSSMAELLCALRRFDVLKVRFGMTRE 80

Score = 60 (27.6 bits), Expect = 7.4e-06, Sum P(2) = 7.4e-06Identities = 12/17 (70%), Positives = 13/17 (76%), Frame = +3

Query: 45 LDTDEKEMLLFLCRDVA 95

LD E E+L FLCRDVA

Sbjct: 18 LDASEHEVLRFLCRDVA 34

Score = 32 (14.7 bits), Expect = 6.8, Sum P(2) = 1.0Identities = 8/16 (50%), Positives = 10/16 (62%), Frame = +3

Query: 162 VGDLAELLYRVRRFDL 209

V L LL+ V R+DL

Sbjct: 148 VSVLVTLLHAVCRYDL 163

ER

pir/1855668 hypothetical protein E8 - equine herpesvirus 2 gi/695247

(U20824) ORF E8 [Equine herpesvirus 2] Length = 171

Plus Strand HSPs:

Score = 66 (30.4 bits), Expect = 4.2e-05, Sum P(2) = 4.2e-05Identities = 15/34 (44%), Positives = 20/34 (58%), Frame = +3 Query: 120 RDLLDILRERGKLSVGDLAELLYRVRRFDLLKRI 221

RD L + LS + ELL+RV R DL++RI

Sbjct: 39 RDAFKFLSDYACLSAANQMELLFRVGRLDLIRRI 72

Score = 55 (25.3 bits), Expect = 4.2e-05, Sum P(2) = 4.2e-05Identities = 10/17 (58%), Positives = 13/17 (76%), Frame = +3

Query: 42 ALDTDEKEMLLFLCRDV 92

+LD DE E L+LCRD+

Sbjct: 12 SLDEDETETYLYLCRDL 28



gil1492102 (U60315) MC159L [Molluscum contagiosum virus subtype 1]

Length = 241

. Plus Strand HSPs:

Score = 63 (29.0 bits), Expect = 9.5e-05, Sum P(2) = 9.5e-05Identities = 15/44 (34%), Positives = 25/44 (56%), Frame = +3

Query: 117 VRDLLDILRERGKLSVGDLAELLYRVRRFDLLKRILKMDRKAVE 248

V L L ++ KL++ L E+LY ++R DLLK + ++ E

Sbjct: 42 VTQALCSLSQQRKLTLAALVEMLYVLQRMDLLKSRFGLSKEGAE 85

Score = 58 (26.7 bits), Expect = 9.5e-05, Sum P(2) = 9.5e-05Identities = 12/28 (42%), Positives = 16/28 (57%), Frame = +3

Query: -12 SAEVIHQVEEALDTDEKEMLLFLCRDVA 95

S + + E LD+ E +LLFLC D A

Sbjct: 9 SLPFLRHLLEELDSHEDSLLLFLCHDAA 36



gil1813378 (D86547) still life type 1 [Drosophila melanogaster]

Length = 2064

Minus Strand HSPs:

Score = 40 (18.4 bits), Expect = 0.00040, Sum P(4) = 0.00040Identities = 10/38 (26%), Positives = 17/38 (44%), Frame = -3

Query: 125 VPDIRWNHIYSNIPAQKEQHLLLICIKCFFNLMDDFSR 12

+PD ++ +Y E+ L C + N M+ F R

Sbjct: 1127 MPDNAYSTVYLRDAMSVEEFLASACARRNLNPMEHFVR 1164

Score = 39 (17.9 bits), Expect = 0.00040, Sum P(4) = 0.00040Identities = 5/11 (45%), Positives = 8/11 (72%), Frame = -3

Query: 299 HQHSIVRNKVR 267

HQH+++ N R

Sbjct: 370 HQHNVINNNTR 380

Score = 35 (16.1 bits), Expect = 0.00040, Sum P(4) = 0.00040 Identities = 8/14 (57%), Positives = 10/14 (71%), Frame = -1

Query: 172 KSPTDSLPLSRKIS 131

+SP DSLP R+ S

Sbjct: 724 RSPWDSLPSLRQDS 737

Score = 33 (15.2 bits), Expect = 0.00040, Sum P(4) = 0.00040 Identities = 7/24 (29%), Positives = 11/24 (45%), Frame = -3

Query: 260 EQVGLHSFSVHLQDTFEQVKSPHS 189

E LH+F L + E + + S

Sbjct: 597 ESYSLHTFEALLSQSMENLANAKS 620

$\mathbf{E} \mathbf{R}$

SPIP47211 GALR HUMAN GALANIN RECEPTOR (GAL1-R) gi | 559048 (L34339)

galanin receptor [Homo sapiens] gi|1297338 (U53511) galanin receptor [Homo sapiens] Length = 349

Plus Strand HSPs:

Score = 73 (33.6 bits), Expect = 0.31, P = 0.26Identities = 12/22 (54%), Positives = 15/22 (68%), Frame = +2

Query: 221 LEDGQKSCGDPPAQEPSPCFGL 286

L +G SC +PPA EP P FG+ jct: 8 LSEGNASCPEPPAPEPGPLFGI 29



gil58271 (X12896) CTAP-III protein (AA 1-86) [Cloning Vector ps29P]

Length = 86

Plus Strand HSPs:

Score = 46 (21.2 bits), Expect = 0.88, Sum P(2) = 0.59 Identities = 11/31 (35%), Positives = 16/31 (51%), Frame = +3

Query: 36 EEALDTDEKEMLLFLCRDVAIDVVPPNVRDL 128

EE+LD+D L LC + P N++ L

Sbjct: 8 EESLDSDLYAELRCLCIKTTSGIHPKNIQSL 38

Score = 42 (19.3 bits), Expect = 0.88, Sum P(2) = 0.59

Identities = 8/13 (61%), Positives = 10/13 (76%), Frame = +2

Query: 221 LEDGQKSCGDPPA 259

L+DG+K C DP A

Sbjct: 56 LKDGRKICLDPDA 68

ER

gil208127 (M88539) Col-CTAP-III (Leu21) small fusion protein [Cloning

vector] gi|208129 (M88538) Col-CTAP-III(Leu21) small fusion protein [Cloning vector] gi|209574 (M88536) connective tissue activating peptide III [Cloning vector]

Length = 91

Plus Strand HSPs:

Score = 46 (21.2 bits), Expect = 1.2, Sum P(2) = 0.70

Identities = 11/31 (35%), Positives = 16/31 (51%), Frame = +3

Query: 36 EEALDTDEKEMLLFLCRDVAIDVVPPNVRDL 128

EE+LD+D L LC + P N++ L
13 EESLDSDLYAELRCLCIKTTSGIHPKNIQSL 43

Score = 42 (19.3 bits), Expect = 1.2, Sum P(2) = 0.70Identities = 8/13 (61%), Positives = 10/13 (76%), Frame = +2

Query: 221 LEDGQKSCGDPPA 259

L+DG+K C DP A

Sbjct: 61 LKDGRKICLDPDA 73



Sbjct:

gil1685110 (U58210) tetrahydrofolate dehydrogenase/cyclohydrolase

[Streptococcus thermophilus]
Length = 284

Minus Strand HSPs:

Score = 68 (31.3 bits), Expect = 1.6, P = 0.79Identities = 13/40 (32%), Positives = 25/40 (62%), Frame = -3

Query: 284 VRNKVRVPEQVGLHSFSVHLQDTFEQVKSPHSVEQFSQVP 165

VRNK R ++ G HS +V+L ++ ++ +E+++Q P

Sbjct: 50 VRNKERAAKKAGFHSKTVNLSESLSEEELIEVIEKYNONP 89



gi|791038 (X84709) mediator of receptor induced toxicity [Homo sapiens]

Length = 208

Plus Strand HSPs:

Score = 51 (23.5 bits), Expect = 2.1, Sum P(2) = 0.87Tidentities = 14/47 (29%), Positives = 22/47 (46%), Frame = +3

Query: 171 LAELLYRVRRFDLLKRILKMDRKAVETHLLRNPHLVSDYRVLMSEIG 311

L ELL +RR DLL+R+ + A L + V+ +G

Sbjct: 63 LRELLASLRRHDLLRRVDDFEAGAAAGAAPGEEDLCAAFNVICDNVG 109

Score = 42 (19.3 bits), Expect = 2.1, Sum P(2) = 0.87Identities = 8/21 (38%), Positives = 12/21 (57%), Frame = +3

Query: 21 VIHQVEEALDTDEKEMLLFLC 83

++H V +L + E L FLC

Sbjct: 7 LLHSVSSSLSSSELTELKFLC 27



pir||A56912 FADD protein - human gi|809487 (U24231) FADD [Homo sapiens]

Length = 208

Plus Strand HSPs:

Score = 51 (23.5 bits), Expect = 2.1, Sum P(2) = 0.87

Identities = 14/47 (29%), Positives = 22/47 (46%), Frame = +3

Query: 171 LAELLYRVRRFDLLKRILKMDRKAVETHLLRNPHLVSDYRVLMSEIG 311

L ELL +RR DLL+R+ + A L + + V+ +G

Sbjct: 63 LRELLASLRRHDLLRRVDDFEAGAAAGAAPGEEDLCAAFNVICDNVG 109

Score = 42 (19.3 bits), Expect = 2.1, Sum P(2) = 0.87

Identities = 8/21 (38%), Positives = 12/21 (57%), Frame = +3.

Query: 21 VIHQVEEALDTDEKEMLLFLC 83

++H V +L + E L FLC

Sbjct: 7 LLHSVSSSLSSSELTELKFLC 27



Sbjct:

gb|1002001 Sequence 1 from Patent US 4897348 gb|110108| Sequence 1 from

Patent WO 8501067 Length = 90

Plus Strand HSPs:

Score = 44 (20.2 bits), Expect = 2.5, Sum P(2) = 0.91Identities = 10/31 (32%), Positives = 16/31 (51%), Frame = +3

Query: 36 EEALDTDEKEMLLFLCRDVAIDVVPPNVRDL 128

EE+LD+D L +C + P N++ L 12 EESLDSDLYAELRCMCIKTTSGIHPKNIQSL 42

Score = 42. (19.3 bits), Expect = 2.5, Sum P(2) = 0.91

Identities = 8/13 (61%), Positives = 10/13 (76%), Frame = +2

Query: 221 LEDGQKSCGDPPA 259

L+DG+K C DP A

Sbjck: 60 LKDGRKICLDPDA 72 '



gil208159 (M11517) connective tissue activating peptide-III [Artificial gene]

Length = 91

Plus Strand HSPs:

Score = 44 (20.2 bits), Expect = 2.6, Sum P(2) = 0.92Identities = 10/31 (32%), Positives = 16/31 (51%), Frame = +3

Query: 36 EEALDTDEKEMLLFLCRDVAIDVVPPNVRDL 128

EE+LD+D L +C + P N++ L

Sbjct: 13 EESLDSDLYAELRCMCIKTTSGIHPKNIQSL 43

Score = 42 (19.3 bits), Expect = 2.6, Sum P(2) = 0.92

Identities = 8/13 (61%), Positives = 10/13 (76%), Frame = +2

Query: 221 LEDGOKSCGDPPA-259

L+DG+K C DP A

Sbjct: 61 LKDGRKICLDPDA 73



splp223661my88 mouse myeloid differentiation primary response protein

MYD88 pir||S11226 MyD88 protein - mouse gi|53294 (X51397) MyD88 protein (AA 1-243) [Mus musculus] Length = 243

Plus Strand HSPs:

Score = 66 (30.4 bits), Expect = 2.9, P = 0.95Identities = 17/42 (40%), Positives = 23/42 (54%), Frame = +3

Query: 96 IDVVPPNVRDLLDILRERGKLSVGDLAELLYRVRRFDLLKRI 221 ++ P R LLD + R SVG L ELL + R D+LK + Sbjct: 11 LETRPDPTRSLLDAWQGRSGASVGRLLELLALLDREDILKEL 52



gil807578 (M19420) unknown protein [Germiston virus]

Length = 75

Plus Strand HSPs:

Score = 43 (19.8 bits), Expect = 3.9, Sum P(2) = 0.98Identities = 5/12 (41%), Positives = 8/12 (66%), Frame = +1

Query: 226 RWTEKLWRPTCS 261

+WT+ WR C+
42 KWTQNFWRRLCA 53

Score = 39 (17.9 bits), Expect = 3.9, Sum P(2) = 0.98Identities = 9/22 (40%), Positives = 12/22 (54%), Frame = +1

Query: 142 GKEVSCLSGTWLNCSTE*GDLT 207 G++ +C S L CS E D T

Sbjct: 8 GQKSTCPSSQVLKCSLELSDST 29

ER

Sbjct:

splP29425|TX26_PHONI_NEUROTOXIN TX2-6 pir||S29216 neurotoxin Tx2 -

spider (Phoneutria nigriventer) gi|256377 neurotoxin Tx2-6 [Phoneutria nigriventer=Brazilian armed spiders, venom, Peptide, 48 aa] Length = 48

Plus Strand HSPs:

Score = 34 (15.6 bits), Expect = 4.3, Sum P(2) = 0.99 Identities = 6/13 (46%), Positives = 8/13 (61%), Frame = +1

Query: 121 GTFWIFYGKEVSC 159

G FWI + K +C

Sbjct: 34 GYFWIAWYKLANC 46

Score = 33 (15.2 bits), Expect = 4.3, Sum P(2) = 0.99Identities = 6/13 (46%), Positives = 6/13 (46%), Frame = +2

Query: 89 CCYRCGST*CQGP 127

CC G C GP

Sbjct: 16 CCGERGECVCGGP 28



splp02775|pF4L HUMAN PLATELET BASIC PROTEIN PRECURSOR (PBP) (CONTAINS:

CONNECTIVE-TISSUE ACTIVATING PEPTIDE III (CTAP-III), LOW-AFFINITY PLATELET FACTOR IV (LA-PF4), BETA-THROMBOGLOBULIN (BETA-TG), NEUTROPHIL-ACTIVATING PEPTIDE 2 (NAP-2)) pir||TGHU beta-thromboglobulin precursor - human gi|181176 (M54995) connective tissue activating peptide III [Homo sapiens] gi|344294 (A01319) novel factor having neutrophil-stimulating activity [unidentified] Length = 128

Plus Strand HSPs:

Score = 44 (20.2 bits), Expect = 8.5, Sum P(2) = 1.0Identities = 10/31 (32%), Positives = 16/31 (51%), Frame = +3

Query: 36 EEALDTDEKEMLLFLCRDVAIDVVPPNVRDL 128
EE+LD+D L +C + P N++ L
Sbjct: 50 EESLDSDLYAELRCMCIKTTSGIHPKNIQSL 80

Score = 42 (19.3 bits), Expect = 8.5, Sum P(2) = 1.0Identities = 8/13 (61%), Positives = 10/13 (76%), Frame = +2

Query: 221 LEDGQKSCGDPPA 259 L+DG+K C DP A Sbjct: 98 LKDGRKICLDPDA 110

Parameters:

V=50 B=50 H=0 -filter=SEG P=4

-ctxfactor=5.61 E=10

: .							
Query				As Used		Com	outed
Frame	. MatID	Matrix n	ame Lambd	a K	н	Lambda I	к н
Std.	0	BLOSUM62				0.319 0.3	135 0.396
.+.3.	0	BLOSUM62	0.319	0.135	0.396	0.320 0.	140 0.377
+2	0	BLOSUM62	0.319	0.135	0.396	0.376 0.3	173 0.727
+1	0	BLOSUM62	0.319	0.135	0.396	0.353 0.3	151 0.594
-1 .	0	BLOSUM62	0.319	0.135	0.396	0.335 0.1	143 0.410
-2	0	BLOSUM62	0.319	0.135	0.396	0.350 0.1	154 0.532
-3	. 0	BLOSUM62	0.319	0.135	0.396	0.334 0.1	143 0.458
. ,					•		
Query		: · · · · .		٠.	٠		• • • • • • • • • • • • • • • • • • • •
Frame	MatID	Length	Eff.Length	E SW	тх	E2 S2	
+3	0	103	103	10. 58 3	13 22	0.11 32	
+2	0.	104	104	10. 58 3	13 22	0.11 32	•
+1	. O ·	104	104	10. 58 3	13 22	0.11 32	.*
· -1	0	104	65	10. 55 3	13 22	0.11 30	
-2	. 0	104	104	10 58 3	13 22	0 11 32	•

58 3

13 22

Statistics:

Query		Expected	Observed	HSPs	HSPs
Frame	MatID	High Score	High Score	Reportable	Reported
+3	0	62 (28.5 bits)	121 (55.7 bits)	18	18
+2	0 .	62 (28.5 bits)	73 (33.6 bits)	7	7
+1	0	62 (28.5 bits)	62 (28.5 bits)	3	· * 3 ·
-1	0	58 (26.7 bits)	50 (23.0 bits)	1	1
-2	0.	62 (28.5 bits)	59 (27.1 bits)	0	0
-3	0	62 (28.5 bits)	68 (31.3 bits)	1 4.	4

Query	Neigl	hborhd Word	Excluded	Failed	Successful	Overlaps
Frame	MatID Word	ds Hits	Hits	Extensions	Extensions	Excluded
+3	045	74 3643889	411910	3212799	19180	7
+2	0 101	18 4815505	897825	3908333	9347	34
+1	0 158	B2 5097249	987386	4099341	10522	8
-1	0 28	37 1981913	251753	1723979	6181	6
-2	0 65	56 4186873	555161	3623591	8121	3
-3	0 94	47 4154885	590204	3549776	14905	7

Database: Non-redundant GenBank CDS translations+PDB+SwissProt+SPupdate Release date:

Posted date: 8:09 AM EST # of letters in database: 68,723,500

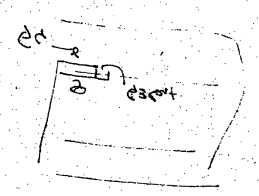
of sequences in database; 241,922 # of database sequences satisfying E: 17

No. of states in DFA: 571 (56 KB) 4 Total size of DFA: 109 KB (128 KB) Time to generate heighborhood: 0.01u 0.01s 0.02t Real: 00:00:00

No. of processors used: 4

Time to search database: 47.57u 0.34s 47.91t Real: 00:00:12

47,64u 0.45s 48.09t Real: 00:00:12



mmest -> List

DNA sequence 313 b.p. agagtaggatgt ... agagattggtga linear

	ı	10	1	20	ı	30		4.0	1	50	1	60	
1	agagtag	gat	gtctgc	tgaa	gtcatc	cate	aggttg	aaga	agcact	tgat	acagat	gaga	60
61	aggagat	gct	gctctt	tttg	tgccgg	gatg	ttgcta						
121	gggacct	tct	ggatat	ttta	cgggaa	agag	gtaage						
181	tgctcta	cag	agtgag	gcga	tttgac	ctgc	tcaaac	gtat	cttoaa	gata	gacaga	aaao	240
	ctgtgga						accttg						
	cagagat			·			•		33	,-	,,,,,,,,	50	313
	ı	10	i	20	1	30		40	1	50	1	60	

Exhibit I

[Image]

[progress meter]
Search in progress, please wait for results

BLASTN 1.4.9MP [26-March-1996] [Build 14:27:07 Apr 1 1996]

Reference: Altschul, Stephen F., Warren Gish, Webb Miller, Eugene W. Myers, and David J. Lipman (1990). Basic local alignment search tool. J. Mol. Biol. 215:403-10.

Notice: this program and its default parameter settings are optimized to find nearly identical sequences rapidly. To identify weak similarities encoded in nucleic acid, use BLASTX, TBLASTN or TBLASTX.

Query= .tmpseq_1 (219 letters)

Database: Non-redundant Database of GenBank EST Division 816,369 sequences; 299,962,672 total letters.

		Smalles	Smallest	
		Sum		
	High	Probabil:	ity	
Sequences producing High-scoring Segment Pairs:	Score	P(N)	N	
gb AA001257 AA001257 zh82c06.rl Soares fetal liver spleen	360	1.2e-63	3	
dbj C05730 C05730 similar to none	250	4.5e-37	3	
gb AA149562 AA149562 zo29e05.rl Stratagene colon (#937204	267	2.1e-33	3	
qb T93307 T93307 ye29c04.sl Homo sapiens cDNA clone 1	191	2.4e-09	2	
gb AA151642 AA151642 zo29e05.s1 Stratagene colon (#937204	204	2.0e-07	1	

Plus Strand HSPs:

Score = 360 (99.5 bits), Expect = 1.2e-63, Sum P(3) = 1.2e-63Identities = 72/72 (100%), Positives = 72/72 (100%), Strand = Plus / Plus

Query: 195 TGGATGGGGCCA 206 |||||||||| Sbjct: 589 TGGATGGGGCCA 600

Score = 353 (97.5 bits), Expect = 1.2e-63, Sum P(3) = 1.2e-63Identities = 71/72 (98%), Positives = 71/72 (98%), Strand = Plus / Plus

```
127 AAAGATGTTTTT 138
 Query:
           111111111111111
 Sbjct:
        520 AAAGATGTNTTT 531
 Score = 213 (58.9 bits), Expect = 1.2e-63, Sum P(3) = 1.2e-63
 Identities = 43/44 (97%), Positives = 43/44 (97%), Strand = Plus / Plus
         25 TCCCAGAGTGTGTATGGTGTGGATCAGACTCACTCAGGGCTCCC 68
 Query:
           417 TCCCAGAGTGTATGGTGTGGATCAGACTCACTCAGGGCTCNC 460
 Sbjct:
 Score = 139 (38.4 bits), Expect = 2.2e-57, Sum P(3) = 2.2e-57
 Identities = 35/45 (77%), Positives = 35/45 (77%), Strand = Plus / Plus
          1 GTGTGTGTCCTGGTGAGCCGAGGATCCCAGAGTGTGTATGGTGTG 45
 Query:
           Sbjct:
        391 GTGTGTGTCCTGGTGAGCCGAGGANNTCCCAGAGTGTGTATGGTG 435
 dbj|C05730|C05730 similar to none
          Length = 521
  Plus Strand HSPs:
- Score = 250 (69.1 bits), Expect = 4.5e-37, Sum P(3) = 4.5e-37
 Identities = 52/55 (94%), Positives = 52/55 (94%), Strand = Plus / Plus
         64 CTCCCTGCATCACATCAGGAGGATGTTCATGGGAGATTCATGCCCTTATCTAGCA 118
Query:
           467 CCCCCTGCATCACATCAGGAAGATGTTCNTGGGAGATTCATGCCCTTATCTAGCA 521
 Sbjct:
 Score = 236 (65.2 bits), Expect = 4.5e-37, Sum P(3) = 4.5e-37
 Identities = 48/49 (97%), Positives = 48/49 (97%), Strand = Plus / Plus
         Query:
           Sbict:
 Score = 120 (33.2 bits), Expect = 4.5e-37, Sum P(3) = 4.5e-37
 Identities = 24/24 (100%), Positives = 24/24 (100%), Strand = Plus / Plus
         1 GTGTGTGTCCTGGTGAGCCGAGGA 24
Query:
           399 GTGTGTGTCCTGGTGAGCCGAGGA 422
Sbict:
gb|AA149562|AA149562 zo29e05.rl Stratagene colon (#937204) Homo sapiens
          cDNA clone 588320 5'
          Length = 538
  Plus Strand HSPs:
 Score = 267 (73.8 bits), Expect = 2.1e-33, Sum P(3) = 2.1e-33
 Identities = 55/57 (96%), Positives = 55/57 (96%), Strand = Plus / Plus
```

Page 2

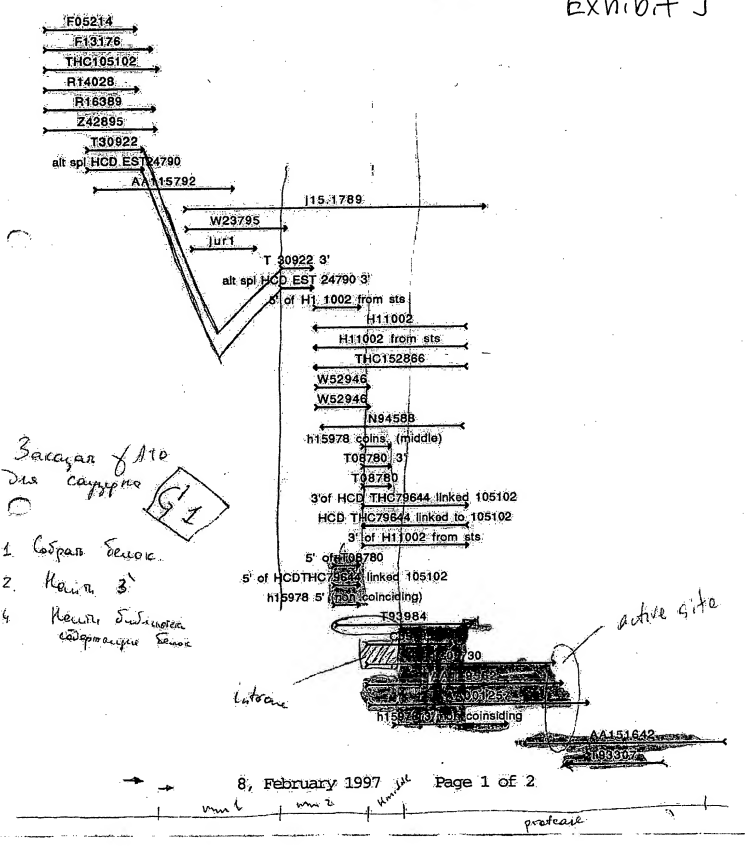
```
66 CCCTGCATCACATCAGGAGGATGTTCATGGGAGATTCATGCCCTTATCTAGCAGGGA 122
Query:
           Sbjct:
        465 CCCTGCATCACATCAGGAGGATGTTCATGGGAGAATCATGCCCTTATCTAGCAGGAA 521
Score = 221 (61.1 bits), Expect = 2.1e-33, Sum P(3) = 2.1e-33
Identities = 45/46 (97%), Positives = 45/46 (97%), Strand = Plus / Plus
         23 GATCCCAGAGTGTGTATGGTGTGGATCAGACTCACTCAGGGCTCCC 68
Query:
           421 GGTCCCAGAGTGTATGGTGTGGATCAGACTCACTCAGGGCTCCC 466
Sbjct:
Score = 135 (37.3 bits), Expect = 1.2e-24, Sum P(3) = 1.2e-24
Identities = 35/45 (77%), Positives = 35/45 (77%), Strand = Plus / Plus
         1 GTGTGTGTCCTGGTGAGCCGAGGATCCCAGAGTGTGTATGGTGTG 45
Query:
           Sbjct:
        397 GTGTGTGTCTGGTGAGCCGAGGAGGTCCCAGAGTGTGTATGGTG 441
Score = 75 (20.7 bits), Expect = 2.1e-33, Sum P(3) = 2.1e-33
Identities = 15/15 (100%), Positives = 15/15 (100%), Strand = Plus / Plus
Query:
        126 CAAAGATGTTTTAT 140
           1111111111111
Sbjct:
        523 CAAAGATGTTTTTAT 537
gb|T93307|T93307 ye29c04.sl Homo sapiens cDNA clone 119142 3'.
          Length = 294
 Minus Strand HSPs:
Score = 191 (52.8 bits), Expect = 2.4e-09, Sum P(2) = 2.4e-09
Identities = 39/40 (97%), Positives = 39/40 (97%), Strand = Minus / Plus
        190 CAAAGAGGCTGCTGTCCTCCAGCTGGCCCTCTGACACCAC 151
Query:
           255 CCAAGAGGCTGCTCTCCAGCTGGCCCTCTGACACCAC 294
Sbjct:
Score = 89 (24.6 bits), Expect = 2.4e-09, Sum P(2) = 2.4e-09
Identities = 21/25 (84%), Positives = 21/25 (84%), Strand = Minus / Plus
       212 TTATCGTGGCCCCATCCACCTCCAA 188
Query:
           234 TCATCGCTGGCCCATCCACCTCCAA 258
Sbjct:
gb|AA151642|AA151642 zo29e05.sl Stratagene colon (#937204) Homo sapiens
          cDNA clone 588320 3' similar to contains Alu repetitive element;
          Length = 608
 Minus Strand HSPs:
Score = 204 (56.4 bits), Expect = 2.0e-07, P = 2.0e-07
Identities = 64/93 (68%), Positives = 64/93 (68%), Strand = Minus / Plus
```

Page 3

```
Query:
        187 AGAGGCTGCTCCTCCAGCTGGCCCTCTGACACCACATAGTTCTGAATAAAAACATCTT 128
                               1111111
        446 AGAGGCTGCTGTCTCAAGCTGGGCCCTCTGACACCACATAGTTCTGAATAAAAAACATCT 505
Sbjct:
Query:
        127 TGGCTTCCCTGCTAGATAAGGGCATGAATCTCC 95
            1 1 1 1 1
                       Sbjct:
        506 TTGGCTTCCCTGCTAGATAAGGGCTGAATCTCC 538
Parameters:
 V=100
 B = 50
 H=0
 P=4
 -ctxfactor=2.00
 E = 10
                            ---- As Used ----
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 Query
                            Lambda K
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                                                   Lambda K
                                                                   H
 Strand MatID Matrix name
                            0.192
                                   0.173
                                           0.357
     0 +5,-4
                                                   same
                                                          same
                                                                  same
          0
             +5,-4
                            0.192 0.173
                                           0.357
  -1
                                                   same
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 Query
                                     s W
                                                    E2 S2
0.021 74
 Strand MatID Length Eff.Length
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                                             T X
             219
                                 10. 111 11
       0
                         219
                                             0 73
          0
                219
                         219
                                 10. 111 11
                                             0 73
                                                    0.021 74
  -1
Statistics:
                                                 HSPs
 Query
               Expected
                               Observed
                                                           HSPs
                              High Score
 Strand MatID High Score
                                              Reportable Reported
     0
              119 (32.9 bits) 365 (100.9 bits)
                                              11
                                                            11
  +1
  -1
              119 (32.9 bits) 204 (56.4 bits)
                                                   3
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              Neighborhd Word
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 Query
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                                                22832
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                                     2286
                                                       1213
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  +1
                          26331
  -1
         0
                 212
                          23923
                                      2040
                                                20821
                                                         1062
                                                                      5
 Database: Non-redundant Database of GenBank EST Division
   Release date:
   Posted date:
 # of letters in database: 299,962,672
 # of sequences in database: 816,369
 # of database sequences satisfying E: 5
 No. of states in DFA: 172 (172 KB)
 Total size of DFA: 178 KB (192 KB)
 Time to generate neighborhood: 0.00u 0.00s 0.00t Real: 00:00:00
 No. of processors used: 4
 Time to search database: 9.20u 0.31s 9.51t Real: 00:00:03
```

Total cpu time: 9.21u 0.32s 9.53t Real: 00:00:03

Exhibit J



for 5' utr sense
5' GTT TCT TTG CCT CCA TCT TGG GTG CGC CTT 3' 18646

fot atg in mm sense
5' ATG TCT GCT GAA GTC ATC CAT CAG GTT GAA 3' 1864*

for before middle reg sense
5' CTG GTT GCC CCA GAT CAA CTG GAT TTA TTA 3' 18648

after middle antisense
5' GCC AAG CTG TTC CTT AAG TCT TTG TTC TTT 3'

first strong homology in prot reg (YKMKSK) sense
5' ACC TGA AGA GAG ATA CAA GAT GAA GAG CAA 3' 18650

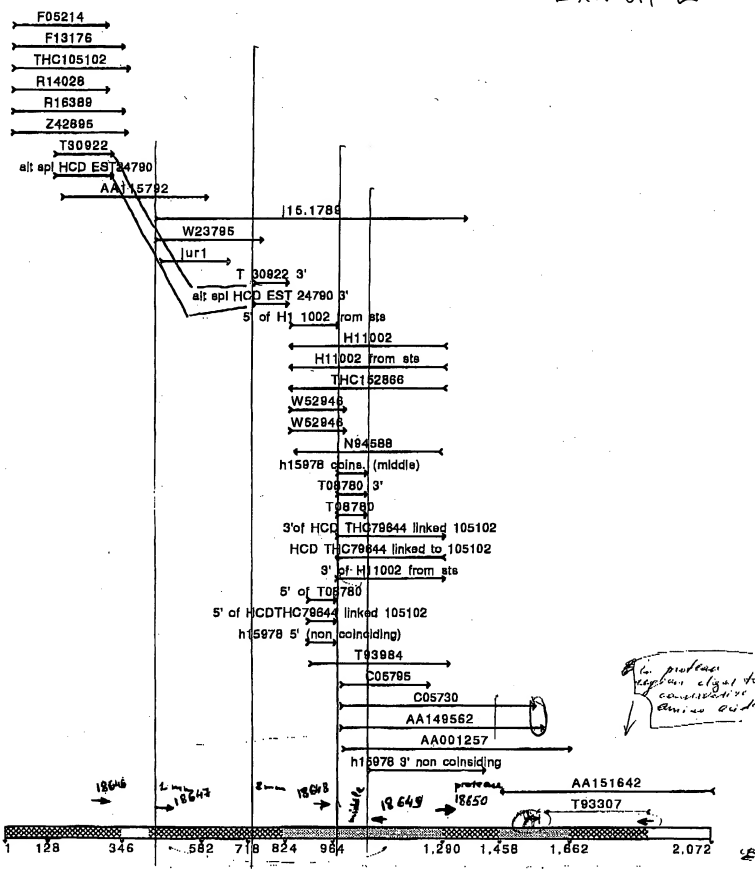
for second subunit for (EADF) antisence
5' CAG TAC ACA GGC TCC AGA AGA AGT CAG CTT 3'

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Exhibit L



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Exhibit M

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[Image]

[progress meter]
Search in progress, please wait for results

TBLASTN 1.4.9MP [26-March-1996] [Build 14:27:13 Apr 1 1996]

Reference: Altschul, Stephen F., Warren Gish, Webb Miller, Eugene W. Myers, and David J. Lipman (1990). Basic local alignment search tool. J. Mol. Biol. 215:403-10.

Notice: statistical significance is estimated under the assumption that the equivalent of one complete reading frame of the database codes for protein and that significant alignments will involve only coding reading frames.

Query= tmpseq_1 (139 letters)

Database: Non-redundant Database of GenBank EST Division 824,500 sequences; 302,742,428 total letters.

			-	Smalles	st
				Sum	
		Reading	High	Probabil	lity
Sequences producing High-scoring	g Segment Pairs:	Frame	Score	P(N)	N
gb AA198928 AA198928 mu23f09.rl	Soares 2NbMT Mus	muscu +3	79	0.040	1
gb AA014850 AA014850 mh28c10.r1	. Soares mouse plac	centa +1	37	0.81	2
gb W23795 W23795 zb79a05.rl	Soares senescent	fibro +3	63	0.9997	1

gb|AA198928|AA198928 mu23f09.rl Soares 2NbMT Mus musculus cDNA clone 640265 5' similar to TR:G695247 G695247 ORF E8.; Length = 421

Plus Strand HSPs:

Score = 79 (36.9 bits), Expect = 0.040, P = 0.040Identities = 25/106 (23%), Positives = 41/106 (38%), Frame = +3

Query: 5 EVLCEVARKLGTDDREVVLFLLNVFIPQPTLAQLIGALRALKEEGRLTFPLLAECXXXXX 64

EV+ +V L D++E++LFL + L +L E G+L+F LAE

Sbjct: 99 EVIHQVEECLDEDEKEMMLFLCRDVTENLAAPNVRDLLDSLSERGQLSFATLAELLYRVR 278

Query: 65 XXXXXXXXXXXXXXXXXERHLAGTMSYFSPYQLTVLHVDGELCARDI 110

E HL S Y++ ++ L D+

Sbjct: 279 RFDLLKRILKTDKATVEDHLRRNPHLVSDYRVLLMEIGESLDQNDV 416

gb|AA014850|AA014850 mh28c10.rl Soares mouse placenta 4NbMP13.5 14.5
 Mus musculus cDNA clone 443826 5'
 Length = 143

Plus Strand HSPs:

Score = 37 (17.3 bits), Expect = 1.7, Sum P(2) = 0.81Identities = 9/14 (64%), Positives = 9/14 (64%), Frame = +1

Query: 87 TMSYFSPYQLTVLH 100

T S F P QLTV H

Sbjct: 19 TCSSF*PSQLTVKH 60

Score = 35 (16.3 bits), Expect = 1.7, Sum P(2) = 0.81. Identities = 5/6 (83%), Positives = 6/6 (100%), Frame = +3

Query: 134 CTVWKT 139

CTVW+T

Sbjct: 63 CTVWQT 80

Length = 313

Plus Strand HSPs:

Score = 63 (29.4 bits), Expect = 8.2, P = 1.0Identities = 18/54 (33%), Positives = 28/54 (51%), Frame = +3

Query: 5 EVLCEVARKLGTDDREVVLFLLNVFIPQPTLAQLIGALRALKEEGRLTFPLLAE 58

EV+ +V L TD++E++LFL + L L+E G+L+ LAE

Sbjct: 18 EVIHQVEEALDTDEKEMLLFLCRDVAIDVVPPNVRDLLDILRERGKLSVGDLAE 179

Parameters:

V=100

B = 50

H=0

-filter=SEG

P=4

-ctxfactor=6.00

E=10

---- As Used Computed Query Н Lambda K Н Lambda K Frame MatID Matrix name 0 0.323 0.138 0.404 same same same BLOSUM62

Query

Frame MatID Length Eff.Length E S W T X E2 S2 +0 0 139 118 10.603 14 22 0.058 34

Statistics:

Query Expected Observed HSPs HSPs Frame MatID High Score High Score Reportable Reported +0 0 63 (29.4 bits) 79 (36.9 bits) 4 4

Neighborhd Word Excluded Failed Successful Overlaps Frame MatID Words Hits Hits Extensions Extensions Excluded +0 0 、489 27788736 3256934 24493093 38709

Database: Non-redundant Database of GenBank EST Division

Release date: February 16, 1997

Posted date: 7:38 AM EST Feb 16, 1997 # of letters in database: 302,742,428 # of sequences in database: 824,500

of database sequences satisfying E: 3

No. of states in DFA: 289 (31 KB) Total size of DFA: 37 KB (64 KB)

Time to generate neighborhood: 0.00u 0.00s 0.00t Real: 00:00:00

No. of processors used: 4
Time to search database: 86.35u 0.35s 86.70t Real: 00:00:22

Total cpu time: 86.37u 0.39s 86.76t Real: 00:00:22

your ID: 10609.1818 ** our Ref: SampleB-1818 16 10609.1818.seq Length: 1021 February 25, 1997 21:08 Type: N Check: 5641

- 1 ACCAAACCAA AAAAAGAGNC CTAGAACTAG TCGGAATTCG GCACGAGGTG
- 51 TCTGCTGAAG TCATCCATCA GGTTGAAGAA GCACTTGATA CAGATGAGAA
- 101 GGAGATGCTG CTCTTTTTGT GCCGGGATGT TGCTATAGAT GTGGTTCCAC
- 151 CTAATGTCAG GGACCTTCTG GATATTTTAC GGGAAAGAGG TAAGCTGTCT
- 201 GTCGGGGACT TGGCTGAACT GCTCTACAGA GTGAGGCGAT TTGACCTGCT
- 251 CAAACGTATC TTGAAGATGG ACAGAAAAGC TGTGGAGACC CACCTGCTCA
- 301 GGAACCCTCA CCTTGTTTCG GACTATAGAG TGCTGATGGC AGAGATTGGT
- 351 GAGGATTTGG ATAAATCTGA TGTGTCCTCA TTAATTTTCC TCATGAAGGA
- 401 TTACATGGGC CGAGGCAAGA TAAGCAAGGA GAAGAGTTTC TTGGACCTTG
- 451 TGGTTGAGTT GGAGAAACTA AATCTGGTTG CCCCAGATCA ACTGGGATTT
- 501 ATTAGAAAAA ATGCCTAAAG AACATCCACA GAATAGACCT GAAGACAAAA
- 551 ATCCAGAAGT ACAAGCAGTC TGTTCAAGGA GCAGGGACAA GTTACAGGGA
- 601 ATGTTCTCCA AGCAGCAATC CAAAAGATCT CAAGGATCCT TCAAATAACT
- 651 TCAGGATGAT AACACCCTAT GCCCATTGTC CTGATCTGAA AATTCTTGGA
- 701 AATTGTTCCa TGTGATTAAC ATGGAACTGC CTCTACTTAA TCNTCTGAAT
- 751 GATAAATCGT TCNTTTTCTA AATGTNTTAT AATGTNTTTA NCCCTTNCTT
- 801 GTTGCGGTAT TTTAAATGCT TCCCATCTTT TGTTACTACT AATATGCNAT
- 851 AAATAAATTC CTTGTNCTCT TAAAAAAAA AAAAAAAAA AAAAAATTCC
- 901 CGCGCCCCA CNCANGGGGG GCCGGTCCCN TCCCNNNTTN ANTTTTTANT
- 951 CCNGGCCCTT TNCCCCTNNT GGGAACCCNT CTTNATCTNA ANGAAACCCC
- 1001 CNTCCTNCNT GCCNCCCCCC C

your ID: 11717.1818 ** our Ref: SampleB-1818 17 11717.1818.seq Length: 1028 February 25, 1997 21:08 Type: N Check: 5259

- 1 CGATTCATAG ATCAGGGTTT TCCCAGTCAC GACGTTGTAA AACGACGGCC
- 51 AGTGAATTGT AATACGACTC ACTATAGGGC GAATTGGGTA CCGGGCCCCC
- 101 CCTCGAGGTC GAGCGGCCGC AGGAATTTTT TTTTTTTTT TTTTTTTTT
- 151 TTTTTTTAA AAAGTACAAG GATATTTATT TTATAGCATT ATTAGTAGTA
- 201 ACAAAAATT GGAAAGCATC TAAACATACA GCAACAANAA AGGGCTAAAC
- 251 ACattataac acattagaa aatgaaacaa titaatcatt caaaatgatt
- 301 AATTAAAGGC AGTtCCATNT TAATCACATG GAACAATTTC CAAAAATTTT
- 351 CAAATCAGGA CAATGGGCAT AGGGTGTTAT CATCCTGAAT TTATTTGAAG
- 401 GATCCTTGAA ACTCTTTTGG ATTGCTGCTT GGAAAACATt CCTGTAACTT
- 451 GTCCCTGCTC CTTGAACAAA CTGCTTGTAC TTCTGGATTT TTGTCTtCAG
- 501 GTCTATTCTG TGGATqTTCT TTAGGCATTT TTCTAATAAA TCCAgTTGAT
- 551 CTGGGGCAAC CAAATTTATT TTCTCCAACT CAACCACAAG GTCCAAAAAA
- 601 CTCTTCTCCT TGCTTATCTT GCCTCGGCCC ATGTTTCCTT CaTGAGGAAA
- 651 ATTAATGAGG ACCTTCANAT TTATCCAANT CCTCNCCAAT CTCTGCCATC
- 701 ACACTCTATA TTCCAAAACA AGGTGANGGT CCTGAACANG TGGGTCTCAC
- 751 ACTITICTGT CCNCTCCAAA TANTTTGAAC AGGTCAAATC CCTCNTCTGT
- 801 TAAACATTCA CCANTTCCCN ACAAAAACTT ACCCTTTCCN TAAAATATCA
- 851 AAGGTCCTGA CTTANGTGAA CCCCTCTATA CCNCTCCCGC CCAAAAAAAC
- 901 CACCCCTCCC NNCTTTCATT CTCTCCCCCG AAGGAAAATC NCAAACCNCT
- 951 CCAATCCANT ATCCAGATCC TTTTNGGGTT GGGGGGTCTC CCCCCAANAA
- 1001 NTTANNNCCC NCCTNTTTGT TTANCCCC

your ID: 18648.1818 ** our Ref: SampleB-1818 18 18648.1818.seq Length: 1012 February 25, 1997 21:08 Type: N Check: 1701 ..

- 1 AANANGCCTA AAGAACAtCC ACAGAATAGA CCTGAAGACA AAAATCCAGA
- 51 AGTACAAqCA GTCTGTTCAA GGAGCAGGGA CAAGTTACAG GAATGTTCTC
- 101 CAAGCAGCAA TCCAAAAGAG TCTCAAGGAT CCTTCAAATA ACTTCAGGAT
- 151 GATAACACCC TATGCCCATT GTCCTGATCT GAAAATTCTT GGAAATTGTT
- 201 CCATGTGATT 'AACATGGAAC TGCCTCTACT TAATCATTCT GAATGATTAA
- 251 ATCGTTTCAT TTTCTAAATG TGTTATAATG TGTTTAGCCC TTTCTTGTTG
- 301 CTGTATGTTT AGATGCTTTC CAATCTTTTG TTACTACTAA TAATGCTATA
- 351 ΑΑΑΤΑΑΑΤΑΤ CCTTGTACTT CTTAAAAAAA AAAAAAAAA AAAAAAAAA
- 401 AAAAATTNCC TGCGGCCGCN CAACCTCAAG GGGGGGCCCG GTACCCAATN
- 451 CGCCCTATAN TNATTCTTAT TACAATNCAC TGGCCGNCGT TTTACAACNT
- 501 CGTNACTGGG AAAAACCCTN ATCTATNAAT CNTAAATACT GAAAAACCCC
- 551 GCAATTTCAC TTCAACTGTG CATCGTGCAC CGTCTCAATT TCTTTCNTTT
- 601 ATACNTCGTT TNGCCTTCTN TTATNTAACT ATACCCCCCN AANTTCCAAT
- 651 CTNGGCCNTG TANCCTCTGA TCTATAAAAA TTTTTAAATG ACTAAAATAA
- 701 NTGCCCCTT TTTTTTGGAC CTNCTTCTCN TGAANTTTNT NACAAAGGCT
- 751 ATCCAAANCT TGGACTCTNC CCNAAGTTGG TCAATCNCNA ACAGGTGTCG
- 801 GNTNTCTACC TTNCCAAAAA TTCCAAAAAA AGNAAAGGTC AANCTTGGTA
- 851 ANCTINITAA ACTCCAATAC CAATCCINNN ANTAINAINN TAINAINAAT
- 901 NATTTAAAAA ANNATNTTAN AATTTAAATT GACCTAGGTT AAACAAAATT
- 951 CAANCEAATA CCNCCGTAGT CNGTCCTNCN GGTAACATAA NCCCTTGACC
- 1001 CCCCACGGCC CC

your ID: rti-651.1831 ** our Ref: SampleC-1831 20 rti-651.1831.seq Length: 765 March 3, 1997 11:26 Type: N Check: 1445 ...

- 1 CTCCGGTGAA CTGTGCACAG CCCTCGCTTC TGAGCCTTGA ATTCCACATT
- 51 CTTCATCGCT GGCCCATCCA CCTCCAAGAG GCTGCTGTTC TCCAGCTGGC
- 101 CCTCTGACAC CACATAGTTC TGAATAAAAA ACaTCTTTGG CTTCCCTGCT
- 151 AGATAAGGGC ATGAATCTCC CATGAACATC CTCCTGATGT GATGCAGGGG
- 201 GAGCCCTGAG TGAGTCTGAT CCACACCATA CACACTCTGG GAGCCTCCTC
- 251 GGCTCACCAG GACACACA AAGCTGTCGT AGTCTCGGTG CTCGGGCATA
- 301 CAGGCAAATT GGCCAAGAAT CTGGGATATA CCATGCATAC TGAGATGCAA
- 351 GAATTTCTGG ACTTCATAGC CCAGGGGAAG TGAAGGTGTC TCGAAGAAGC
- 401 TCTGTCTCAT TGCCAATGCA ATCGATTATC AGGCAGATTC CTAGGGGCTT
- 451 GCTCTTCATC TTGTATCTCT CTTCAGGTAT GCTCTGAGGC-AAAAAAGCTT
- 501 CTGATTCCtG AATGGATTTC TTCACTGGTT CTTGTtGAGC GCCAAGCTGT
- 551 TCCTTAAGTC TTTGTTCTTT ACTTCTCCCA TTATGGAGCC TGAAGTTATT
- 601 TGAAGGATCC TTGAGACTCT TTTGGATTGC TGCTTGGAAA ACATTCCTGT
- 651 AACTTGTCCC TGCTCCTTGA ACAGATGCTT GTATTCTNGG ANTTTTGTCT
- 701 CAGGTCTANT CTGTGGATGT CTTTAGGCAT TTTCTCATAA ATCCACTTGA
- 751 TTTGGGCACA AATTT

your ID: rt1-648.1831 ** our Ref: SampleC-1831 19 rt1-648.1831.seq Length: 763 March 3, 1997 11:26 Type: N Check: 6251 ...

- 1 AAAATGCTÀA AGAaCaTCCA CAGAATAGAC CTGAAGACAA AAATCCAGAA
- 51 GTACAAGCAG TCTGTTCAAG GAGCAGGGCA CAAGTTACAG GAAATGTTCT
- 101 CCAAGCAGCA ATCCAAAAGA GTCTCAAGGA TCCTTCAAAT AACTTCAGGC
- 151 TCCATAATGG GAGAAGTAAA GAACAAAGAC TTAAGGAACA GCTTGGCGCT
- 201 CAACAAGAAC CAGTGAAGAA ATCCATTCAG GAATCAGAAG CTTTTTTGCC
- 251 TCAGAGCATA CCTGAAGAGA GATACAAGAT GAAGAGCAAG CCCCTAGGAA
- 301 TCTGCCTGAT AATCGATTGC ATTGGCAATG AGACAGAGCT TCTTCGAGAC
- 351 ACCTTCACTT CCCTGGGCTA TGAAGTCCAG AAATTCTTGC ATCTCAGTAT
- 401 GCATGGTATA TCCCAGATTC TTGGCCAATT TGCCTGTATG CCCGAGCACC
- 451 GAGACTACGA CAGCTTTGTG TGTGTCCTGG TGAGCCGAGG AGGCTCCCAG
- 501 AGTGTGTATG GTGTGGATCA GACTCACTCA GGGCTCCCCC TGCATCACAT
- 551 CAGGAGGATG TTCATGGGAG ATTCATGCCC TTATCTAGCA GGGAAGCCAA
- 601 AGATGTTTTT TATTCAGAAC TATGTGGTGT CAGAAGGCCA gCTGGAgAAC
- 651 AGCACCTCTT GGAGGTGGAT GGCCACCANN AAAATGTGGA ATTCCAGGCT
- 701 CACAACCAAG GCTGTTGCCA TTTCACGACA AACTGANTTC CTCTGGAACC
- 751 TGTGTANGCG GGA

your ID: v1-t7.1822 ** our Ref: SampleB-1822 23 v1-t7.1822.seq Length: 775 February 25, 1997 21:05 Type: N Check: 6815 ...

- 1 TCNTTGGGCG CGCTTGTCGA CTGGCCGAGG CGGCCTTTTT TTTTTTTTT
- 101 GTACAAGGAT ATTTATTTTA TAGCATTATT AGTAGTAACA AAAGATTGGA
- 151 AAGCATCTAA ACATACAGCA ACAAGAAAGG GCTAAACACA TTATAACACA
- 201 TTTAGAAAAT GAAACGATTT AATCATTCAA AATGATTAAG TAAAGGCAGT
- 251 TCCATGTTAA TCaCATGGAA CAATTTCCAA AAATTTTCAG ATCAGGACAA
- 301 TGGGCATAGG GTGTTATCAT CCTGAAGTTA TTTGAAGGAT CCTTGAAACT
- 351 CTTTTGGATT GCTGCTTGGA AAACATTCCT GTAACTTGTC CCTGCTCCTT
- 401 GAACAACTG CTTGTACTTC TGGATTTTTG TCTTCAGGTC TATTCTGTGG
- 451 ATGTTCTTTA GGCATTTTTC TAATAAATCC AGTTGATCTG GGGCAACCAA
- 501 ATTTAGTTTC TCCAACTCAA CCACAAGGTC CAAGAAACTC TTCTCCTTGC
- 551 TTATCTTGCC TCGGCCCATG TAATCCTTCN TGANGAAAAT TAATGAGGAC
- 601 CCTCCNATTT ATCCAAATCC TCCCCAATCT CTGCCATCAN CACTCTATAN
- 651 TCCGAAACAA GGTGANGGTT CCTGAACAGG TGGGTCTCCC ANCTTTTCTG
- 701 TCCATCTTCC AAANACNTTT GAACAGGTCA ATCCCCCNCC CGTTAAACAT
- 751 TCCACCAATT CCNANAAAAC TTACT

your ID: v1.1822 ** our Ref: SampleB-1822 17 v1.1822.seg Length: 821 February 25, 1997 21:05 Type: N Check: 3460 ..

1 GGANCGTCGA GGCATTACAA TCGCGAAACC AAGCCATAGC ATGAAACAGC 51 GAGCTTGCAG CCTCACCGAC GAGTCTCAAC TAAAAGGGAC TCCCGGAGCT 101 AGGGGTGGGG ACTCGGCCTC ACACAGTGAG TGCCGGCTAT TGGACTTTTG TCCAgTGACA GCTGAGACAA CAAGGACCAC GGGAGGAGGT GTAGGAGAGA 151 AGCGCCGCGA ACAGCGATCG CCCAGCACCA AGTCCGCTTC CAGGCTTTCG 201 GTTTCTTTGC CTCCATCTTG GGTGCGCCTT CCCGGCGTCT AgGGGAGCGA 251 301 ACGCTGAGGT GGCAGCGGCa GGAGAGTCCG GCCGCGACAG GACTAACTCC CCCaCTGGAA ACGATTCTGA AAGAAATGAA GTCAGCCCTC ATAAATGAAG 351 TTGACTGCCT GCTGGCTTTC TGTTGACTGG CCCGGAGCTG TACTGCaAGA 401 CCCTNGTGAG CTTCCCTAgT CTAAGAGTAN GATGTCTGCT GAAGTCaTCC 451 ATCAGGTTGA aCAagCCNTG ATNCCGATCA NANNGAGATC CTGCCCTTTC 501 551 TGTGCCGGGG ANGTTNCTAT ANATGTGGTT CCNCCTTANT NTCTGGGACC 601 NTCTGGTATT TTTCACNGNN AACAGGTNCC TNTCTGTCCN GGAATNNGCT 651 NACTNCTCCA CTCAATNANG CGATNTNACC TGCTCNACTT CCTNCNATAT 701 GGACTNNCAT CCTGTGGAAN NCCTCTTNCT CATGACCCCT CTCCTTTTNC 751 CGTACTNACT ATTCNCCNGC TCAANATTNN TTAAGNNCTG NAANANTCNA

801 TCTTNTTCAT TATTTTTGCT C

your ID: v2-t7.1822 ** our Ref: SampleB-1822 24 v2-t7.1822.seq Length: 820 February 25, 1997 21:05 Type: N Check: 4660 ...

- 1 AAAGAGATGG GATCTTGCTA TATTGCCCAG GTTGGTATGC AGTGGCTATt
- 51 CGTAGGCACA ATCACAGCAT GCTATGACCC TGAACTCCTG GGTTCNNAAg
- 101 ATCCTCCCAC CTCAGCCTCC CaAGTAACTG GGAATNCAGG TACCCACACC
- 151 CACACCCANC TAATTTTTGC ATTTCTACTA GGGACAGCGT TTACCATGTT
- 201 GGCCAGGCTG GTCTCNAACt CCTGACCTGA AGTGATCTGC CCTCCTTGGC
- 251 CTCCCAAAGT GCTGGGATTA CAGGCGTGAG CCACTACaCC CACCCTTTTG
- 301 GTTTCTTATG TGTAGGAqAq GATAAGTTTC TTTCTCAGAa TGTGCTGCaq
- 351 CCAGACATAA TATTTCTCCT TGGCAGAAAC TNTGCNGTTC CAATCATACA
- 401 TGTAGCCATT GCANTTCAAT GTGAANATCC ANGAAGTGGG CGTTTTCTTT
- 451 CNAGTCTCNN TTTCTGGGAN ANGCACTNCN NGTACCGGAC NGTGAGCTGN
- 501 TGAAACTGCT CCATCAGGGA ANTGTNTGCA TTACACACGG CTCCATAAAN
- 551 AAATCNNCTT CCNCGNTGAA AATGTGCACN GCCCTCGCTN CTTNATCCTT
- 601 GAAATNNNTC TTCCTCATCC TNNGNCCNTC CCCNCNAAAA TGGTGCTGTC
- 651 TCCCAACTGG CCCCGAAAC ACATTTTCTC NNAATNTCNA NNCNNANGGC
- 701 TCCCTGCCA TATGGCTNAA NCGCCCGAAC TCCCCTGAAC TAACCTGGGG
- 751 ACCNCCTNA TTNGNAACTN CCACCCCNCC CTNGTACTCT TTGNTCCCTC
- 801 AATTCATNAC CTTTTANTCC

your ID: v4-t7.1822 ** our Ref: SampleB-1822 25 v4-t7.1822.seq Length: 797 February 25, 1997 21:05 Type: N Check: 6426 ..

- 1 TCNTTGNGCG CGCTTGTCGA CTGGCCGAgG CGGCCTTTTT TTTTTTTTT
- 101 TTTTTTTTT TTTTTTTTT AAAAATTNCA AGGATNTTTA TTTTATANCA
- 151 TTATTAGTAT TAACAAAAA TTGGAAAGCA TCTAAACATA CAGCAACAAA
- 201 AANGGGNTAA ACNCNTTATA ACNCNTTTAA AAAATGAAAC AATTTAATCN
- 251 TTCAAAATGA TTAATTNAAG GCAGTNCCAT GTTAATCNCN TGGAACAATT
- 301 TCCAAAAATT TTCAAATNAG GACAATGGGC ATAGGGTNTT NTCNNCCTGA
- 351 ATTTATTTGA AGGATCCTTG AAACNCTTTT GGATTGCTGC TTGGAAAACA
- 401 TTCCTGTAAC TTGTCCCTGC NCCTTGAACA AACTGCTTGT NCTTCNGGAT
- 451 TTTTGTCTNC AGGTCTATTC TGTGGATNTT CTTTAGGCAT TTTTCTAATA
- 501 AATCCNNTTN ATCTGGGGCA ACCAAATTTA TTTTCTCCAA CTCAACCACA
- 551 AGGTCCAAAA AACTCTTCTC CTTGCTTATC TTGCCTCGGC CCATGTTATC
- 601 CTTCCTGAGG AAAATTAATG AGGACCCTCN AATTTATCCA AATCCCCCCA
- 651 ATCTCTGCCA TCANCACTCT ATANTCCGGG TTTCCCNAAT GGTAATGGCC
- 701 TCAAANGATC CNTATCNANA TGTCCCNAAA TNATCCGGCA AAAAAAAACC
- 751 CGGGCGNTTC NGTCCCAACG AAANCAAAAT TCNAACGGAG GCGGACG

your ID: v4.1822 ** our Ref: SampleB-1822 19 v4.1822.seq Length: 836 February 25, 1997 21:05 Type: N Check: 7807 ...

- 1 GGACCGTCGA GGCCATTACA ATCGCGAAAC CGGACTATAG AGTGCTGATG
- 51 GCAGAGATTG GTGAGGATTT GGATAAATCT GATGTGTCCT CATTAATTTT
- 101 CCTCATGAAg GATTACATGG GCCGAGGCAA GATAAGCAAG GAGAAGAGTT
- 151 TCTTGGACCT TGTGGTTGAG TTGGAGAAAC TAAATCTGGT TGCCCCACAT
- 201 CAACTGGATT TATTAGAAAA ATGCCTACAG AACATCCACA CAATNCACCT
- 251 GAAGACCAAA ATCCAGAAGT ACAAGCATTC TGTTCAAGGA GCAGGGACAA
- 301 GTTACCGAA TGTTCTCCNN TCANCNATCC AAAAGANTCT CANNGGATCC
- 351 TTCACATAAC TTCNCGATGA TNACACCCTA NGCCCNTTGT CCTGATCTGA
- 401 AAANTCTCGG AANTTGTTCC ANGTGATTAA CATNGAACTG CCNCTNCNNA
- 451 NTCANTCTGA ATGACTAAAT CGTTTCACNT CCTNAATGTG TTANANTGTG
- 501 TNTANCCCCT TCNTGTTGCC TGTATGTCTA CATNCTTTCC NNTCNTTCGT
- 551 TACTNCTANT AATGCTCTNN TATATNTATC CTTGTNCTTC TCTCTCTTCC
- 601 NCTCCCNCTT CTTCNTCTCC TTTCCTCCCT CCCNCCNTNC TCCCTTCTCT
- 651 TNCCTCTCTC TCNNNNCATT AAGGCNCCCC GCTACTTCAA NGGCCCCCAN
- 701 GGNCCCCNAC ANTINITCTT NTTTTTACCC TCCCCCTTNC CCCNNAAGTT
- 751 GCCCGGTAAN TCCGNCTCCT TANTTTTCTN CCCCCCTTCC CCTCNCCCAT
- 801 CGGAANACTT NNTTATTTCG TCTCTCANNC NCNNNC

your ID: v5-t7.1822 ** our Ref: SampleB-1822 26 v5-t7.1822.seq Length: 793 February 25, 1997 21:05 Type: N Check: 9524 ...

- 1 AANTACAAGG ATATTTATTT TATAGCATTA TTAGTAGTAA CAAAAGATTG
- 51 GAAAGCATCT AAACATACAG CAACAAGAAA GGGCTAAACA CATTATAACA
- 101 CATTTAGAAA ATGAAACNAT TTAATCATTC AGAATGATTA AGTNNAGGCA
- 151 GTTCCATGTT AATCACATGG AACAATTTCC AAAAATTTTC AGATCAGGAC
- 201 AATGGGCATA GGGTGTTATC ATCCTGAAGT TATTTGAAGG ATCCTTGAAA
- 251 CTCTTTTGGA TTGCTGCTTG GAAAACATTC CTGTAACTTG TCCCTGCTCC
- 301 TTGAACAGAC TGCTTGTACT TCTGGATTTT TGTCTTCAGG TCTATTCTGT
- 351 GGATGTTCTT TAGGCATTTT TCTAATAAAT CCAGTTGATC TGGGGCCAACC
- 401 AGATTTAGTT TCTCCAACTC AACCACAAGG TCCAAGAAAA CTCTTCTCCT
- 451 TGCTTATCTT GCCTCGGCCC ATGTAATCCT TCATGAGGAA AATTAATGAG
- 501 GACACATCAG ATTTATCCAA ATCCTCACCA ATCTCTGCCA TCAGCACTCT
- 551 ATAATCCGAA ACAANGTGAA GGTTCNTGAA CACGTGGGTC TCCACACCTT
- 601 TTCTGTCCTC TTCAAAATAC TTTGAACAGG TCAATCCCCa CTCTGTAAAC
- 651 ATTCACCCAG TCCCGACNAN NCTTACTCTT TCCNTAAATN CCNAANGTCC
- 701 NACTTAGTNG AACNCCCTTT NCTCCCCCGN CCAAAAANCC CCCTCNTCGT
- 751 TTAATNNTCN CCCCGANGAG ACNTCCANTC NNCTNACAGG NAC

your ID: v6-t7.1822 ** our Ref: SampleB-1822 27 v6-t7.1822.seq Length: 779 February 25, 1997 21:05 Type: N Check: 1979 ...

- 1 GAAGTACAAG GATATTTATT TTATAGCATT ATTAGTAGTA ACAAAAGATT
- 51 GGAAAGCATC TAAACATACA GCAACAAGAA AGGGCTAAAC ACATTATAAC
- 101 ACATTTAGAA AATGAAACGA TTTAATCATT CAGAATGATT AAGTAGAGGC
- 151 AGTTCCATGT TAATCACATG GAACAATTTC CAAGAATTTT CAGATCAGGA
- 201 CAATGGGCAT AGGGTGTTAT CATCCTGAAG TTATTTGAAG GATCCTTGAG
- 251 ACTCTTTTGG ATTGCTGCTT GGAqAACATT CCTGTAACTT GTCCCTGCTC
- 301 CTTGAACAGA CTGCTTGTAC TTCTGGATTT TTGTCTTCAG GTCTATTCTG
- 351 TGGATGTTCT TTAGGCATTT TTCTAATAAA TCCAGTTGAT CTGGGGCAAC
- 401 CAGATTTAGT TTCTCCAACT CAACCACAAG GTCCAAGAAA ACTCTTCTCC
- 451 TTGCTTATCT TGCCTCGGCC CATGTTATCC TTCATGAGGA AAATTAATGA
- 501 GGACACATCA GATTTATCCA AATCCTCACC AATCTCTGCC ATCAGCACTC
- 551 TATAGTCCGA AACAAGGTGA AGGTTCCTGA ACAGGTGGGT CTCCACAGCT
- 601 TTTCTGTCCA CTTCAAANAC TTTTGAACAG GTCAATCNCC TCACTCTGTT
- 651 AAACATTCAC CATTTCCCGA CAAANNCTTA CTCTTTCCCG TNAAATACCN
- 701 AAAGGTCCTG ACATTTGTGG AACTCNTCTA TTACACNTCC CGGCNCAAAA
- 751 AACACNCCNC TTCTCTNNNT TCAATTTCT

